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### **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: C07K 15/06, 15/08, 15/12 C12N 15/13, C12P 21/08 G01N 33/563

(11) International Publication Number:

WO 91/06570

A1 | (4

(43) International Publication Date:

16 May 1991 (16.05.91)

(21) International Application Number:

PCT/AU90/00513

(22) International Filing Date:

25 October 1990 (25.10.90)

(30) Priority data:

PJ 7045

25 October 1989 (25.10.89) AU

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(81) Designated States: AT, AT (European patent), AU, BB, BE (European patent), BF (OAPI patent), BG, BJ (OAPI patent), BR, CA, CF (OAPI patent), CG (OAPI patent), CH, CH (European patent), CM (OAPI patent), DE, DE (European patent), DK, DK (European patent), ES, ES (European patent), FI, FR (European patent), GA (OAPI patent), GB, GB (European patent), GR, GR (European patent), HU, IT (European patent), JP, KP, KR, LK, LU, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL, NL (European patent), NO, RO, SD, SE, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent), US.

#### **Published**

With international search report.

(54) Title: HYBRID Fc RECEPTOR MOLECULES

#### (57) Abstract

A novel immunoglobulin binding molecule is provided being a hybrid of two or more binding molecules such as to provide a hybrid immunoglobulin binding molecule having chimeric properties characteristic of two or more individual immunoglobulin binding molecules in one molecule. Methods of production and applications of such hybrid molecules is also provided.

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# TITLE: Hybrid Fc Receptor Molecules

This invention relates to immunoglobulin binding molecules and in particular to Fc receptor (FcR) molecules and hybrids or chimeric forms thereof capable of binding one or a plurality of classes of antibody.

The invention also encompasses the different forms of Fc receptor molecules including soluble, unbound and bound forms as protein molecules and nucleotide sequences coding therefore, methods of production of hybrid Fc receptor molecules, uses of hybrid Fc receptor molecules including diagnostic testing and pharmaceutical application, poly and monoclonal antibodies detecting hybrid Fc receptor molecules and the uses thereof.

To date the Fc receptor molecules and cDNA clones thereof have been limited to specific antibody class targets for example IgG, IgE, etc.

Receptors for immunoglobulin (Fc receptors or FcR) play key roles in the immune response in allergy and in resistance to infection (1-3). On phagocytes they are responsible for the binding and removal of immune complexes. On T and B cells they are involved in signalling and in the regulation of antibody synthesis (1,2). Secreted Fc receptor related molecules (immunoglobulin binding factors) have been defined that also regulate antibody synthesis and membrane bound or soluble FcR play a role in T cell function (2). The role of FcR in allergy has also been elucidated and the binding of IgE to its receptor plays a pivotal role in these conditions (3).

In addition, to this wide range of function, FcR receptors for all immunoglobulin classes have been defined. In man and the mouse two classes of receptors for IgG have been defined and are designated FcYRI, FcYRII. In addition a third class of receptor FcYRIII has been identified in man. These receptors can be distinguished by (i) structural differences, (ii) affinity of binding of antibody, (iii) specificity for Ig classes or subclasses and (iv) reaction with monoclonal antibodies to FcR. Thus, FcYRI is a high affinity receptor for monomeric IgG whereas FcYRII and FcYRIII bind monomeric IgG with low affinity but bind immune complexes with high avidity (1-2).

Molecular cloning of cDNA or genes encoding these receptors has demonstrated that they are homologous proteins that have evolved to perform their

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unique functions (4,5,6,7). Indeed these cloning experiments have shown that not only are there multiple forms of each of FcyRI, FcyRII in man and mouse (and FcyRIII in man) but that FcyRs are related (4-10). Furthermore, the receptors for IgG and IgE are also related at the nucleotide and protein levels (5,6,11,12).

The interaction of immunoglobulins with FcR was investigated by constructing chimeric FcyR. Mouse FcyRI and FcyRII cDNAs have been cloned and encode related but distinct proteins. FcyRI and II are both typical membrane spanning proteins (4,5,7). The extracellular region of FcyRI is organised into three disulphide bonded domains. By contrast, the extracellular region of FcyRII is composed of only two disulphide bonded domains. Domains 1 and 2 (D1 and D2) of FcyRII are homologous to domains 1 and 2 (d1 and d2) of FcyRII showing approximately 47% amino acid homology and D3 of FcyRI is unique.

In addition to the differences in structure between FcyRI and FcyRII these receptors differ in their specificity and affinity for IgG. Fc RI binds only mouse IgG2a with high affinity whereas FcyRII binds mouse IgGI, IgG2a and IgG2b.

Thus in vitro mutagenesis was used to determine functional regions of these FcyR and to generate chimeric FcyR with the combined properties of these receptors.

In addition in vitro mutagenesis was performed to generate hybrid FcR composed of functional regions of FcyRII and FceRI wherein such a hybrid RcR possess the properties of both FcyRII and FceRI.

Accordingly the invention provides a hybrid FcR molecule capable of binding to any one or more classes of antibody molecules.

Preferably the hybrid FcR comprises one or more functional domains of FcR linked to the structural or functional region of another molecule.

Preferably the hybrid FcR molecule binds to one of antibody classes IgM, IgG, IgA, IgD or IgE.

Preferably the hybrid FcR molecule is in soluble form.

Preferably the hybrid FcR molecule contains domains D1 and D2 of FcyRI linked to the transmembrane region and cytoplasmic tail of FcyRII and is designated FcyRI/II.

Preferably the hybrid FcR molecule contains domain d1 and d2 of FcyRII

linked to domain D3, transmembrane region and the cytoplasmic tail of Fc RI and is designated FcyRII/I.

Preferably the hybrid FcyR is human FcR.

Preferably the hybrid FCYR is mouse FCR.

Preferably the IgE binding region of IgE FcR is linked or located within FcyRI.

Preferably the IgE binding region of IgE FceR is linked or located within FcyRII.

The invention further provides a hybrid FcR wherein at least one functional region comprises an extracellular domain, or parts thereof, derived from FcyRII having the following amino acid sequence:—

- Phe Ser Arg Leu Asp Pro Thr Phe Ser Tle Pro Gln Ala Asn His Ser His Ser Gly Asp Tyr His Cys Thr Gly Asn Ile Gly Tyr Thr Leu Phe Ser Ser Lys Pro Val Thr Ile Thr Val, or functional equivalent thereof.

The invention further provides a hybrid FcR wherein at least one functional region comprises an extracellular domain, or parts thereof, derived from FceRI having the following amino acid sequence: -

- Trp Leu Leu Gin Ala Ser Ala Glu Val Val Met Glu Gly Gln Pro Leu Phe Leu Arg Cys His Gly Trp Arg Asn Trp Asp Val Tyr Lys Val Ile Tyr Tyr Lys Asp Gly Glu Ala Leu Lys, or functional equivalent thereof. Preferably the FcR is soluble or membrane bound.

The invention further provides:

A hybrid FcR being a chimera comprising a first extracellular domain of FcγRII, a second extracellular domain of FcεRI and a transmembrane region and cytoplasmic tail of FcγRII;

A hybrid FcR being a chimera comprising a first extracellular domains of FceRI containing amino acids 1 - 169 of FceRI linked to amino acids 170-281 of FcyRII;

A hybrid FcR being a chimera comprising a first extracellular domain of FceRI being amino acids 1-86 linked to a second extracellular of FcyRII being amino acids 87-169 linked to a transmembrane region and cytoplasmic tail of FcyRII being

amino acids 170-281;

A hybrid FcR being a chimera comprising a first extracellular domain of FceRI containing amino acids 1-86 linked to part of the second extracellular domain of FceRI being amino acids 87-128 linked to part of the second extracellular domain of FcyRII being amino acids 129-169 linked to a transmembrane region and cytoplasmic tail of FcyRII being amino acids 170-281;

A hybrid FcR being a chimera comprising a first extracellular domain of FceRI being amino acids 1-86 linked to part of the second extracellular domain of FcyRII being amino acids 87-128 linked to part of the second extracellular domain of FeRI being amino acids 129-169 linked to a transmembrane and cytoplasmic tail of FcyRII being amino acids 170-281;

A hybrid FcR being a chimera comprising a first extracellular domain of Fc RII being amino acids 1-86 linked to part of the second extracellular domain of Fc RII being amino acids 87-128 linked to part of the second extracellular domain of Fc RII being amino acids 129-169 linked to the transmembrane and cytoplasmic tail of Fc RII being amino acids 170-281;

A hybrid FcR being a chimera comprising the first extracellular domain of FcγRII being amino acids 1-86 linked to the second extracellular domain of FcεRI being amino acids 87-169 linked to a transmembrane and cytoplasmic tail of FcγRII being amino acids 170-281;

A hybrid FcR being a chimaera comprising the first extracellular domain of FcγRII being amino acids 1-86 linked to part of the second extracellular domain of FcεRI being amino acids 87-128 linked to part of the second extracellular domain of FcγRII being amino acids 129-169 linked to the transmembrane and cytoplasmic tail of Fc RII being amino acids 170-281.

The invention further provides amino acid and nucleotide sequences for hybrid FcR, polyclonal and monoclonal antibodies raised thereto, biological products incorporating FcR hybrids, diagnostic kits and assays and pharmaceuticals.

The invention further provides a method of generating hybrid FcR molecules comprising linking different functional domains of different Fc receptor molecules to construct hybrid FcR molecules.

The invention further provides a method of generating hybrid FcR molecules comprising ligating a first FcR molecule at the transmembrane junction, attaching a second FcR molecule to excise suitable domain regions and linking these two fragments to construct a hybrid Fc receptor molecule.

To generate these receptors an Apa-I restriction site was introduced into the cDNA sequence of FcyRI at the junction of sequences encoding D2 and D3. FcyRII cDNA already has an Apa site at the junction of d2 and the membrane spanning region. Thus the introduction of this restriction site (or any convenient restriction site) would facilitate the exchange of sequences between these or other receptors.

To introduce the Apa I site into the FcyRI cDNA two separate PCR mutagenesis reactions were performed and each used oligonucleotide primers containing an Apa I recognition sequence (Fig. 1) (13–15). The template for PCR was the FcyRI cDNA cloned into the pGEXII expression vector (7). PCR amplification of sequences FcyRI D1 and D2 was primed using a oligonucleotide containing Sal I site that hybridized to the leader sequence at the 5' end of the cDNA and the second primer MDHI which hybridized between D2 and D3 and would introduce an Apa I site between nucleotides 591 and 592 by altering three nucleotides of the FcyRI sequence GAGCTC to GGGCCC (Fig. 1). The second PCR amplified the sequence containing D3, the transmembrane domain and cytoplasmic tail. A primer (MDH2) also containing an Apa I recognition sequence hybridizes to the sequences between D2 and D3 of FcyRI in combination with the second oligonucleotide (MDH4) which contained a Sal I site. This primer hybridizes to the pGEXII vector sequences at the 3' end of the FcyRI cDNA insert.

The PCR products were then digested with Apa I or Sal I and ligated into FcyRII expression systems as outlined below.

To construct expressable FcR chimeric cDNA, Fc RII cDNA was subcloned into the PstI site of the expression vectors pKC3 or pKC4 downstream of the SV40 early promoter. These vectors differ only in the orientation of the polylinker and provided a unique Sal I site at the 5' or 3' end of the FcyRII cDNA (Fig. 2a). The pKC3-FcRII and pKC4-FcRII cDNA were digested with Sal I and Apa I to remove FcyRII d1 and d2 or the transmembrane and cytoplasmic encoding regions from these

vectors (Fig. 2b, 2c). The PCR products containing Sal I and Apa I sticky ends and encoding D1 and D2 of Fc RI (generated above; (Fig. 1)) were then subcloned into Sal I/Apa I digested pKC3-FcγRII. Similarly, to generate a hybrid cDNA encoding d1 and d2 of FcγRII linked to D3, and transmembrane and cytoplasmic tail of FcγRI. The PCR products containing D3, the transmembrane and cytoplasmic regions of FcγRI were linked to the D1 and D2 of FcγRII in the pKC4-Fc RII (Fig. 2e).

The chimeric cDNA were checked for the correct sequence by nucleotide sequencing. As expected the FcyRI/II chimera contained FcyRI nucleotide sequence from positions 9 to 594 that encodes FcyRI D1 and D2 and FcyRII cDNA sequence from positions 595 to 1244 that encode the transmembrane and cytoplasmic region of FcyRII (Fig. 3a). The FcyRII/I chimera contained FcyRII nucleotide sequence from positions 1-662 (encoding d1 and d2) and FcyRI sequence from positions 663-1348 that encode D3, the transmembrane and cytoplasmic tail of FcyRI (Fig. 3b).

The specificity of the chimeric FcyR and FcyRI and FcyRII were then tested in a transient expression system by transfecting the cDNA clones into monkey COS cells. The binding of rabbit IgG and mouse Ig subclasses was then assessed by EA rosetting. As expected rabbit IgG bound to cells transfected with FcyRI (Figure 4a) and FcyRII (Figure 4b) and also bound to both chimeric FcyR i.e. FcyRI/II (Figure 4c) and FcyRII/I (Fig. 4d, Table 1). Thus the D1 and D2 of FcyRI (now anchored in the cell membrane by the membrane spanning region of FcyRII) have the capacity to bind IgG in the absence of D3 of FcyRI.

Similarly, d1 and d2 of FcyRII retain the Ig binding capacity when associated with domain 3 of FcyRI (Fig. 1b, Table 1). Since FcyRI and FcyRII differ in their specificity for mouse IgG subclasses (FcyRI binds IgG2a but FcyRII binds IgGI, IgG2a, IgG2b (ref 1–10)) it was of interest to determine the specificity of the chimeric FcyR molecules for mouse IgG subclasses. As expected cells transfected with the native FcyRI bound IgG2a sensitised erythrocytes (Figure 4f) but did not bind IgGI (Figure 4g) and cells transfected with vector only did not bind IgG-EA of any class (Figure 4h, Table 1). The FcyRI/II chimaera (with the extracellular region consisting only of DI and D2 of FcyRI) not only bound IgG2a (Figure 4i) but also

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bound IgGI (Figure 4j) and IgG2b (Table 1), which do not bind to native FcyRI (Table 1; refs 1-10) i.e. these FcyRI domains have now lost their unique specificity for IgG2a and IgG2b and have acquired the specificity of FcyRII. Thus, the specific binding of IgG2a by normal FcyRI involved D1 and D2 as well as D3 since the removal of D3 (by generation of the chimeric FcyRI/II) results in a broadening of the specificity of these domains. Thus by engineering the cDNA we were able to generate receptors of altered specificity i.e. convert FcyRI to FcyRII-like function.

Examination of the chimeric FcyR (FcyRII/I) containing d1 and d2 of FcyRII linked to D3, the transmembrane and cytoplasmic region of FcyRI, showed that this molecule bound IgGI, IgG2a and IgG2b, i.e. the specificity of d1 and d2 of FcyRII was retained. Furthermore, the presence of D3 of FcyRI did not influence the binding of the various mouse IgG subclasses to this chimeric FcyR.

The chimeric FcR proteins expressed on the surface of transfected cells were tested for reactivity with the 2.4G2 monoclonal antibody which reacts with FcyRII but not with FcyRI.

It is known that this antibody completely inhibits Ig binding to FcyRII and does not bind to FcyRI.

Transfected cells expressing normal and chimeric FcγR were pretreated with Fab fragments of the monoclonal antibody 2.4G2 and then tested for their ability to bind rabbit IgG coated erythrocytes. The Fab fragments completely inhibited binding of antibody coated erythrocytes to FcγRII and the chimeric FcγRII/I (Table 2). In addition, the 2.4G2 Fab<sub>2</sub> did not inhibit binding to FcγRI but by contrast completely inhibited binding to the chimeric FcγRI/ II containing D1 and D2 FcγRI (Table 2). Thus D1 and D2 of FcγRI now express the 2.4G2 when not associated with D3 of FcγRI.

To further define the role of the FcyRI extracellular domains, the binding of monomeric IgG to the chimaeric receptors was investigated. The affinity of monomeric IgG binding was assessed by Scatchard analysis with <sup>125</sup>I labelled monoclonal antibodies of IgG2a subtype on transfected COS cells expressing either of the two chimaeric receptors, native FcyRI or FcyRII. The high affinity receptor FcyRI bound IgG2a with affinity 1.4x10<sup>8</sup>M<sup>-1</sup> (Figure 5) which is similar to that

reported previously (1-3,7). In contrast, the binding of monomeric IgG2a by the FcyRI/II and FcyRII/I chimaeric receptors, and the low affinity receptor FcyRII (as expected), was not detectable (not shown).

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Furthermore, high affinity binding was not able to be generated by linking D3 of FcyRI to d1 and d2 of FcyRII, despite the high homology of FcyRII d1-d2 with FcyRI D1-D2. This suggests that all three extracellular domains of FcyRI are required for the high affinity interaction with IgG2a, D3 being crucial in modifying the D1-D2 interaction with IgG binding properties of FcyRII, as they do not detectably bind monomeric IgG2a and also show the broader specificity for IgG1, 2a, 2b immune complexes that is characteristic of FcyRII. Clearly these domains represent a conserved FcyRII-like IgG binding unit between these otherwise functionally distinct receptors. However, there must be specialised differences between the homologous two domain structure of FcyRI and FcyRII, as only D1 and D2 of FcyRI, not d1 and d2 of FcyRII, have the capacity to interact with D3 to produce the specific high affinity binding of IgG2a.

It should be noted that the strategy used to generate the FcyRII/I chimaeric cDNA led to the replacement of the glutamic acid and leucine residues at positions 172 and 173 with glycine and proline, through the introduction of an Apal site at the junction of D2 and D3 of FcyRI. To ensure that these amino acid substitutions were not responsible for the observed properties of the FcyRII/I chimaeric receptor, mutagenesis was used to replace the glycine and proline residues with the original glutamic acid and leucine residues. This chimaeric receptor exhibited identical IgG binding characteristics to the original FcyRII/I chimaera (data not shown).

Based on findings presented herein, it is possible to make several statements on the origin of FcyRI which highlight important aspects regarding evolution of FcyR genes and the Ig superfamily. Given that there is considerable amino acid homology between D1-D2 of FcyRI and d1-d2 of FcyRII(7), and that D1-D2 adopts the specificity and affinity for IgG of FcyRII after the removal of D3, it is clear that the two receptors have a common evolutionary history. Indeed, the recent cloning and mapping of multiple mouse and human FcyRII genes, multiple human FcyRI genes and FcyRIII genes, and both the mouse and human FceRI genes, indicate there has

been considerable gene duplication in the same region of chromosome 1 to produce this subfamily of Ig related molecules which show great amino acid homology (16– 19).

The substantial amino acid identity between the extracellular regions of IgG receptors, (approximately 90% within a class e.g. FcyRIIa as compared to FcyRIIb1, FcyRIIb2 and FcyRIIb3 or FcyRIII-1 compared to FcyRIII-2) or between the extracellular regions between FcR classes (e.g. approximately 50% between FcyRI and FcyRIII and 40% between FcyRII and FceRI) indicates shared structure although there are clear differences in function or specificity (4-12). This high conservation of amino acid identity between FcR is also seen between the low affinity IgG FcyRII and the high affinity IgE receptor FceRI. Although these receptors have very distinct specificity for immunoglobulins, FcyRII binds IgG whereas FceRI binds IgE, these FcR share approximately 40% amino acid identity in their Ig binding, extracellular domains.

Thus it should be possible to generate many novel chimaeric Fc receptors that contain parts and therefore characteristics and functions of multiple receptor classes e.g. a chimaeric FcR composed of FcyRII and FceRI sequences that binds IgG and IgE; a chimaeric FcyR composed of parts of FcyRII and FcyRI or FcyRII and FcyRIII, or FcaR (IgA receptors) and FcyRII or other FcR. Similarly these chimeric receptors may be composed of sequences dervied from three or more Fc receptors e.g. FcyRII and FcyRI and FceRI or FcyRI and FcaR and FceRI and FcuR (IgM receptors) etc.

The origin of the third domain in the extracellular region of FcyRI is also of interest, unlike FcyRII and FcyRIII, FcyRI is unique in that it has an additional extracellular domain. Whether D3 of FcyRI arose by duplication of a related exon or by a insertion of an exogenous exon is not clear, as sequence comparisons indicate D3 is only distantly related to D1 and D2, or to domains of other Ig superfmaily members. However the data presented herein demonstrates how Ig-superfamily genes may have evolved, in that FcyRI probably originated through the acquisition of an exon encoding an intact Ig like domain (D3), presumably by a primordial FcyRII-like gene, with subsequent divergence refining the interaction of the three

domains to produce a new receptor with modified IgG binding characteristics i.e. a specialised high affinity receptor for IgG2a. This model for FcyRI evolution is consistant with the proposed evolution of the Ig superfamily postulated by Williams and colleagues, wherein primordial genes acquire individual exons encoding Iglike domains, and these domains become functionally incorporated to creat new molecules with new function. Indeed, each of the extracellular Ig-like domains of mouse FcyRI and FcyRII are encoded by individual exons (N. Osman, M. Hogarth unpublished observations, Ref. 19).

Finally, based on the high homology between D1, D2 and D3 of human and mouse FcyRI (69,74 and 75% amino acid identity respectively (7), together with the observation that human FcyRI also binds mouse IgG2a with high affinity (10), it is likely that the contributions of the extracellular domains of human FcyRI to IgG binding will parallel those reported here for mouse FcyRI and FcyRII.

Clearly it is possible to generate functional chimeric FcR and several statements can be made. First chimeric cDNA clones encoding FcR composed of components of different FcyR have been generated. Second, by connecting D1 and D2 of FcyRI to the transmembrane cytoplasmic regions of FcyRII a receptor molecule was produced that has a broader specificity than the receptor from which the Ig binding regions were derived, i.e. FcyRI/II containing D1 and D2 of FcyRI binds mouse IgGI, IgG2a, IgG2b. Thirdly, similar experiments that generate chimeric FcyR between human FcyRII and human FcyRI have shown that these chimeric receptors have altered specificity for IgG. Fourthly, it is possible to generate numerous other chimeric FcR that possess characteristics of other FcR. Since most FcR are homologous proteins it is possible to insert/or attach Ig binding regions to FcyRI or II e.g. insert the IgE binding region of IgE FcR (FceRI) into another FcR e.g. FcyRII.

Indeed, the following experiments describe the production of chimaeric receptors containing amino acid sequence elements of the human low affinity FcyRll and the human high affinity IgE receptor FceR1. These chimeras were generated by exchanging parts of the cDNA sequence encoding the regions shown in the Figures 6,7 and Tables 3,4. Although the IgG and IgE receptors have distinct biological

properties and specificites they are homologous transmembrane glycoproteins, both containing two extracellular immunoglobulin binding domains (EC domains) which have approximately 40% amino acid identity. This considerable identity presumably indicates that the extracellular portions of these distinct molecules would be expected to have a similar general structure. Chimeric receptors formed between these classes of receptors would therefore be expected to be expressed on the cell surface, processed appropriately and indeed the transfection experiments indicate that this is the case (Figure 8, Table 5). We have successfully isolated and transfected the human FcyRII cDNA defined in Australian Patent Serial No. 595623 (reference 6) and the cDNA sequence is shown in Figure 6. The human high affinity receptor for IgE has been previously described (12) and the sequence is shown in Figure 7. This receptor is expressed extensively on mast cells and basophils and shows a high affinity for IgE. Biochemical studies have indicated that the Ig binding subunit (a subunit) (sequence shown in Figure 7) is associated with two additional subunits, the β subunit and a dimer of two γ subunits that are disulphide bonded to one another. Expression of the  $\alpha$  subunit normally requires the expression of one or more of the  $\beta$  or  $\gamma$  subunits (21). The strategy employed to examine the Ig binding characteristics of the extracellular domains of FcyRII and FceRI was to exchange the entire domains or parts of domains between two types of receptors and examine the specifity of these receptors with respect to the binding of IgG and IgE. The nucleotide sequence of all chimeric cDNA described herein (Table 3,4) was confirmed by nucleotide sequencing.

Since the FceRI subunit requires the presence of additional subunits for expression and the subunits are not synthesised in the COS cells (which are used as the target cells for transfection and expression experiments) an FceRI  $\alpha$  chain capable of being expressed alone was generated. The initial strategy involved placing the entire extracellular domains of FceRI on the membrane spanning and cytoplasmic region of FcyRII. This was performed by splice overlap extension (SOE) using oligonucleotides EG6 and EG1 (Table 4) which hybridise to the FceRI cDNA (Figure 7) to generate amplified FceRI sequence and the oligonucleotide pair EG2 and EG5 (Table 4) which hybridise to the FcyRII cDNA (Figure 6) to generate the amplified

FcyRII sequence. Since sequence within EG1 and EG2 overlap, the two amplified products were used in the overlap extension to produce the full length recombinant Chimaera 1 (Table 3). Thus a chimaeric cDNA was generated which encoded the 25 amino acids of the FceRI leader sequence and 169 amino acids of the extracellular portion of FceRI which was linked to the amino acid sequences from position 170 in FcyRII to the C terminal amino acid as position 281 of FcyRII (Table 3 and 4).

To establish if this chimeric cDNA encoded a functional IgE receptor the chimeric cDNA contained in expression vector pKC3 (Van Doren et al., J. Virol. 50:606, 1984) was transfected into COS cells in a transient expression system and expression measured 48–72 hours later. Expression was tested by EA rosetting using IgE sensitised erythrocytes and by a direct binding assay using Scatchard analysis using 125 iodine labelled human IgE. Cells transfected with the Chimaera 1 cDNA bound IgE immune complexes, [IgE sensitized erythrocytes (Figures 8a)] and also bound monomeric human IgE with high affinity, but did not bind IgG sensitised erythrocytes (Figures 8d, Table 5) as would be expected of an FceRI. Conversely, FcyRII bound IgG-EA (Figure 8b) but not IgGE-EA (Figure 8A) as expected. This experiment establishes that the expression of Chimaera 1 and the interaction with and specificity for IgE of the Ec domains of FceRI are not dependent on the additional subunits or the membrane spanning region of FceRI.

In the light of experiments using FcR, with the high and low affinity IgG receptors (described earlier) a series of chimeric cDNAs and Fc receptor proteins containing FcyRII and FceRI sequences were then produced in order to:-

- 1. Establish the extent to which replacement of extracellular sequences from one receptor with those of a second receptor would enable the production of functional Fc receptors.
- 2. Generate functional receptors that have properties of multiple Fc receptors e.g. Fc receptors that bound both IgG and IgE.
- 3. Localise the regions of Fc receptors repsonsible for the interaction with immunoglobulins.

To determine the functional Fc binding domain(s) of of FceRI and FcyRII, chimaeric receptors were generated such that they contained two extracellular

domains (EC domains), one of which was derived from FceRI, the second of which was derived from Fc RII (Table 3,4,5). Chimera 2 contained the first extracellular domain of FcyRII and the second domain of FceRI organised in that order. This chimaera was generated by splice overlap extension (SOE) using the oligonucleotides NR1 and EG11 as well as EG10 and EG5 to respectively amplify the appropriate sequences from FcyRII cDNA or from the Chimaera 1 cDNA (Table 4, Figures 6,7). The overlap extension reactions were the performed as detailed in the Materials and Methods and were possible because of the overlapping sequence contained in oligonucleotides EG10 and EG11. The resulting chimaeric cDNA (Chimaera 2) encoded a chimaeric Fc receptor containing the FcyRII leader sequence (amino acid-34 to the N-terminus) and the first EC domain of FcyRII amino acid 1 to 86 inclusive, constituting the first EC domain of the Chimera 2. In addition the overlap extension reactions also included the second EC domain of FCeRI corresponding to amino acids 87 to 169 of this receptor. As Chimaera 1 was used as a template for this region, Chimaera 2 also contains the membrane spanning region and cytoplasmic tail of FcyRII, which included amino acids 170 to 281 (Table 3). Similarly, the chimaeric Fc receptor (Chimaera 3) was generated to contain the first EC domain of FceRI and the second EC domain, transmembrane region and cytoplasmic tail of FcyRII. Splice overlap extension was used to generate the chimeric cDNA that encoded this receptor by using oligonucleotide pair EG6 and EG9 on the FceRI cDNA template and oligonucleotides EG8 and EG5 with FcyRII cDNA as a template. The splice overlap extension reactions were possible because of the overlapping sequence contained in oligonucleotides EG9 and EG8. The Fc receptor encoded by this chiaeric cDNA contained the leader sequence (amino acids-25 to -1 of FceRI) as well as the first EC domain of FceRI (amino acids 1 to 86) as well as the amino acids of the second EC domain, membrane spanning and cytoplasmic tail of FcyRII (amino acids 87 to 281 inclusive) (Tables 3,4 Figures 6,7).

To assess the specificity of these chimaeric receptors, the chimaeric cDNAs, cloned into the pKC3 expression vector, were transiently expressed by transfection into COS 7 cells. The capacity of these Fc receptors expressed on the transfected cells to bind IgG or IgE was tested by the binding of immune complexes in the form

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of erthyrocytes sensitised with IgG of IgE (IgG-EA or IgGE-EA). Chimaera 2 containing the first EC domain of FcyRII and the second EC domain of FCɛRI bound IgE but did not bind IgG sensitised erthrocytes (Figure 8e,f; Table 5). Conversely, Chimaera 3 containing the first EC domain of FcɛRI and the second EC domain of FcyRII bound IgG but not IgE immune complexes (Figure 8g,h). These results indicate that the second EC domain is intimately involved in the interaction with immunoglobulins since in Chimaera 2 the first EC domain was derived from the IgG receptor and the second from the IgE receptor but only bound IgE not IgG. Conversely, in Chimaera 3 the first Ec domain was derived from the IgE receptor and the second from the IgG receptor and could clearly bind IgG but not IgE. Since the normal Fc RII and normal FceRI receptors exclusively bind IgG and IgE respectively (as does Chimaera 1) it is clear that the interaction of Chimearas 2 and 3 with IgE and IgG respectively is mediated within the second EC domain of Fc receptors (Table 3).

To localise the binding of immunoglobulins to a subregions of EC domain 2 in both FcyRII and FceRII additional chimaeric receptors were generated wherein EC domain 1 was derived from either FceRI and FcyRII (Table 4). These chimeric receptors were generated using the splice overlap extension technique, where amplification of the appropriate regions of cDNA were performed on the wild type FCERI or FCYRII cDNA as well as the appropriate chimaeric cDNA (Table 3, 4). Chimaera 4 contained the first EC domain of FceRI together with a substantial amino terminal portion of the second EC domain of FceRI (amino acid -25 to -1 together with animo acids 1 to 128 inclusive). These amino acids were connected to the Cterminal portion of the second EC domain of FcyRII, and membrane/ cytoplasmic regions of FcyRII amino acids 129 to 281 inclusive (Table 3). This chimaeric cDNA was generated using the oligonucleotides EG6 with EG14 on the FceRI cDNA template, and EG15 together with EG5 on the FcyRII cDNA template (Table 4). COS cells were transfected with this chimaeric cDNA and the binding of IgE and IgG to the encoded Chimaera 4 receptor was examined by the binding of immune complexes in the form of antibody sensitized erythrocytes (Figure 8i, j; Table 5). These experiments indicated that the chimaeric Fc receptor Chimaera 4 bound both

IgG-EA and IgE-EA, this indicated that an IgG binding site could be localised to the C-terminal portion of the second EC domain within the region encoded by amino acids 129 to 169 inclusive. The observation that Chimaeria 4 bound IgE-Ea also indicates that an IgE binding region is located in the second domain in the region containing amino acids 87 to 128 inclusive. The involvement of these regions in the binding of IgE and IgG was verified by the contruction of Chimera 5 and its subsequent transfection and immune complex binding studies.

Chimera 5 was generated by the splice overlap extension techniques using the oligonucleotides outlined in Table 4. This chimaera contained the first extracellular domain of FceRI with the second EC domain composed of both FcyRII (amino acids 87 to 128 inclusive and FceRI sequence (amino acids 129 to 169). Transfection of this cDNA indicated that the chimaeric Fc receptor failed to bind IgG or IgE (Tables 3 and 5). The amino acid sequence of Chimeras 4 and 5 differs only in the amino acid sequence within the second EC domain. Wherein amino acid residues, 87 to 128 inclusive, of Chimaera 4 have been derived from FceRI and have been replaced with residues 87 to 128 of FcyRII in Chimaera 5. Since Chimaera 4 binds IgE and Chimaera 5 does not, residues (87-128) in FceRI are important in IgE binding. Similarly as Chimaera 4 contains amino acid residues 129 to 169 inclusive, derived from FcyRII and binds IgG but Chimaera 5 which contains amino acids 129 to 169 inclusive, derived from FceRI and does not bind IgG, indicates that these residues are important in IgG binding. Since Chimaeras 4 and 5 are half domain chimaeras ie. containing sequence from FceRI and FcyRII within the second domain but have the same first EC domain derived from FceRI, two additional chimaeras were made again containing the same amino acid configurations of the second EC domain as found in chimaeras 4 and 5, but containing the first EC domain derived from FcyRII. Chimaera 6 contains the first EC domain of FcyRII including the leader sequence corresponding to amino acids -34 to 1 and amino acids 1-86 inclusive as found in the first EC domain of FcyRII (Table 3, Figure 6). This was attached to a second EC domain wherein amino acids 87 to 128 inclusive were derived from FceRI and amino acids 129 to 169 inclusive were derived from FcyRII (Figures 6.7). Transfection of this cDNA into COS cells showed that the encoded chimaeric

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receptor could bind IgG and IgE (Table 3). By contrast, Chimaera 7 which contained the EC domain 1 of FcγRII but contained amino acids 87–128 of FcγRII and amino acids 129 to 169 of FcεRII but contained amino acids 87–128 of FcγRII and amino acids 129 to 169 of FcεRI failed to bind IgG or IgE when transfected into COS cells (Table 5). These results taken together with thos experiments using the Chimeras 4 and 5 indicate that amino acids within the second domain are clearly important in the interaction of Fc receptors with Ig, and particularly amino acids 87–128 of FcεRI are important in the interaction with IgE and amino acids 129 to 169 of FcγRII in the interaction with IgG.

To further characterise the chimaeric receptor structures, monoclonal anti FcγRII antibodies (description of antibodies described below) were used in immunofluorescence studies on cells transfected with native FcγRII or chimaeric cDNAs. All monoclonal anti FcγRII antibodies bound to cells transfected with the native FcγRII cDNA (Table 5). As expected none of these antibodies bound to cells transfected with the native FcεRI α subunit or cells transfected with Chimaera 1, which contains the extracellular IgE binding domains (Table 5).

Analysis of monoclonal antibody (MAb) binding to the single domain chimaeras indicated that Chimaeras 2 and 3 have some obvious structural differences (Table 5). Monoclonal antibodies 8.7 and 7.30 detected the expression of Chimaera 3 but did not bind to Chimaera 2. Since Chimaera 3 contains EC domain 1 from FceRI and EC domain 2 from FcyRII, this result indicates that the epitope on FcyRII molecules detected by 8.7 and 7.30 (see below) is located in Ec domain of FcyRII. By contrast, monoclonal antibodies 8.2 and 8.26 which bound to the native FcyRII failed to bind to any of the chimaeric receptors tested (Table 5), indicating that this epitope detected by these antibodies requires the presence of part or all the first and second EC domains and this clearly establishes the epitope detected by 8.2 and 8.26 as distinct from the epitope detected by antibodies 8.7 and 7.30. Of interest was the finding that the previously described (22) monoclonal antibody IV.3 bound to Chimaera 3, whereas a second previously described antibody CIKM5 (23) did not bind to any chimaeric cDNA. However, both antibodies bound to the native FcyRII.

The construction of these chimaeric receptors and the immune complex

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binding studies as well as the characterisation using monoclonal antibodies has indicated several clear features of these receptors:-

- The EC domains of FcεRI can be attached to the membrane spanning and cytoplasmic regions of a different molecule (FcγRII) and the binding of IgE to the extracellular domains of FcεRI is retained i.e. the extracellular domains function as receptors irrespective of additional sequence added at the C-terminal end of the EC domains. Clearly soluble forms of this receptor i.e. a form of the receptor containing only the extracellular domains or part thereof in the absence of a membrane spanning segment or cytoplasmic tail, would be expected to bind IgE. Furthermore, the chimaeric receptors which contained appropriate FcεRI or FcγRII sequences (e.g. as found in Chimaeras 4 and 5) would also be expected in a soluble form to bind IgG and/or IgE.
- The second EC domain, EC domain 2, is intimately involved in the binding of IgE and IgG. Indeed, the binding of these immunoglobulins to Chimera 4 and Chimera 6 indicate that the amino acids that correspond to residues 87 to 128 inclusive of FcεRI are intimately involved with IgE binding and residues corresponding to amino acid residues 129 169 of FcγRII are intimately involved in IgG binding (Tables 3–5, Figure 8). These results identify these residues of the domain 2 as being important in the binding of IgE and IgG but do not exclude other regions of the receptor also being intimately involved.
- It is clearly possible to construct multi-functional chimaeric Fc receptors that have properties of several different receptor types. Such multi-functional receptors would be expected to have advantages over individual receptor classes in that the production of a single chimaeric form of Fc receptor would have the functions of multiple Fc receptor classes and would circumvent the need for the production of two individual Fc receptor types with subsequent mixing of these receptors in any pharmaceutical preparation or assay system. The uses of these chimaeric receptors in the diagnosis and treatment of allergy, autoimmune disease, parasite infections, immune complex disease and a range of haemopoietic and non-haemopoietic disorders would be

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significantly more straight forward, than using single function receptors.

Thus the manipulation of FcR will enable the construction of novel chimaeric FcR that can be high or low affinity and bind immunoglobulin of multiple classes. Receptors will have domain or subdomain sequences derived from different receptors (e.g. Ig binding domain or sequences of FcεRI or high affinity domains (D3) of FcγRII that are intimately involved in a particular functions resulting in a single receptor type having multiple functions).

In addition, these functional regions could be attached to non FcR molecules e.g. antibody molecules, or bacterial proteins e.g. to create fusion proteins, wherein FcR function is maintained in addition to the function of the non FcR protein.

5) Furthermore, the monoclonal antibody experiments indicated that the EC domain structure is sufficiently conserved in some case eg. Chimaera 3, and not only interacts with Ig but retains the epitope detected by the monoclonal antibodies 8.7 and 7.30. However, some alteration to the total overall structure is apparent since the epitope detected by antibody 8.2 and 8.26 has been lost from Chimaera 3 as a result of the construction of the chimaeric receptors containing sequences from the FceRI.

The studies described below of the characterisation of these monoclonal anti FcyR antibodies identify antibodies 8.2, 8.26, 8.7, 7.30 as unique monoclonal antibody which are embodied in this patent application.

The characterisation of the native and chimaeric FcR was aided by the use of monoclonal antibodies (MAb). These antibodies recognise cell surface FcyR and also soluble circulating FcR. This has been made possible by the production and characterisation of 4 new MAb (designated 8.2, 8.7, 8.26, 7.30) recognising human FcyRII is described and the use of these MAb in the detection of soluble FcyRII (sFcyRII) in a two antibody radioimmunoassay.

Four MAbs were derived by cell fusion for characterisation. Three MAb [8.2 (of the IgG1 subclass), 8.7 (IgG1) and 8.26 (IgG2b)] were produced by immunisation of BALB/c mice with K562 cells. A fourth "second generation" MAb 7.30 (IgG1) was produced by immunisation with FcyRII 3.0 cDNA transfected L-cells. Cell fusions were performed as described in the Materials and Methods.

#### Tissue Distribution by FACS Analysis.

The tissue distribution of the antigens detected by the MAb was determined by quantitative analysis of fluorescence staining of FcyRII 3.0 transfected L-cells (Tf3.0) and a number of haemopoietic cell lines. This analysis shows that the four new MAb have strong specific positive reactions with FcyRII 3.0 cDNA transfected L-cells, similar to the two CDW32 reference MAb, IV.3 and CIKM5 described by others (22, 23) which are included for the purposes of comparison. The tissue distribution of the new MAbs on the haemopoietic cell lines have profiles which distinguish these from each other and from IV.3 and CIKM5. All antibodies bound strongly to K562 cells but had unique binding on Daudi cells where only MAb 8.7 and 7.30 bound strongly to Daudi (Table 6). The MAb 8.26 bound less strongly and antibody 8.2 showed very weak binding as did the IV.3 and CIKM5 antibodies. Weak binding to Raji cells was also obtained with MAb 8.7, 8.26 and 7.30 and there was no detectable binding with the 8.2 antibody. All antibodies bound to an EBV transformed B cell line (Table 6). It should be noted that the MAb 8.26 gave consistently higher fluorescence or B cell lines (Daudi, Raji, EBV cells) than did the 8.2 and IV.3 MAb.

The binding to myeloid cells was also examined. Antibodies bound to the U937, Thp-1 and HL60 cell lines. However 8.7 and 8.26 could be distinguished from other MAb as they bound less strongly to U937 cells (Table 6). The fluorescence profile s of the binding of the MAb to Daudi and U937 cells is shown in Figure 9. Antibodies IV.3, CIKM5, 8.2 and 8.26 react with greater fluorescence intensity on U937 than on Daudi. Conversely antibodies 8.7 and 7.30 exhibit greater fluorescence than 8.2 or 8.26 or IV.3 and CIKM5. It should be noted that 8.26 has greater reactivity on all B cell lines and can be distinguished from 8.2, IV.3 and CIKM5 (Figure 9). All 6 MAb were negative on the 4 T-cell lines tested.

# Inhibition of Fc Binding.

To determine the relationship of the epitopes detected by these antibodies to the Ig binding site, the capacity of these antibodies to inhibit immune complex (IgG-EA) binding was assessed (Table 7). Results of EA inhibition assays on three FcyRII

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bearing cell lines is shown in Table 2. The three cell lines K562, FcyRII cDNA transfectants (Tf3.0) and Daudi cells express only FcyRII. The pattern of inhibition separates the 6 MAb into 3 groups; IV.3, 8.2 and 8.26 give 80–100% inhibition of EA rosetting on all 3 cell lines. The 8.7 and 7.30 antibodies completely inhibit EA rosetting on Daudi cells, but only partially inhibited EA rosetting on K562 and transfected L-cells. CIKM5 alone, gives partial inhibition of EA on K562 only. Competitive Inhibition Assays.

To Determine Relationship Between Epitopes detected monoclonal antibodies. Competitive inhibition assays between the MAb were initially performed on K562 cells (Table 8) and on transfectants expressing FcyRII (Figure 10). When tested on the K562 cells radiolabelled IV.3, 8.2 and 8.26 show similar complete reciprocal inhibition patterns with unlabelled IV.3, 8.2 and 8.26 i.e. they completely inhibit each others binding indicating these MAb recognise overlapping epitopes. The epitope detected by the IV.3 antibody was unique, as unlabelled IV.3 (but not 8.2 and 8.26) completely inhibited the binding of labelled 8.7, 7.30 and CIKM5. Antibodies 8.7 and 7.30 formed a second distinct group as these completely inhibited each others binding to K562 cells indicating these detect identical epitopes. However, 8.2 or 8.26 did not inhibit the binding of 8.7 and 7.30. Unlabelled CIKM5 either failed to inhibit or only partially inhibited the binding of all other anti-FcyR MAbs tested (Table 8). As discussed below, these results taken together with the tissue distributions and EA inhibition studies indicate the presence of at least 4 epitopes:- Epitope 1 detected by the IV.3 antibody. Epitope 2 detected by the CIKM5 antibody. Epitope 3 by the 8.2 and 8.26 antibodies and Epitope 4 detected by antibodies 7.30 and 8.7.

The competitive binding experiments were also performed on FcyRII cDNA transfected L-cells which express a single form of FcyRII (Figure 10). One MAb defining each of the 4 possible epitopes was studied (IV.3, CIKM5, 8.2, 7.30). The results of competitive inhibition assays are shown in Figure 10. The studies with the Fc RII transfected cells confirms the pattern of inhibition seen when K562 cells were used as target cells.

#### Immunoprecipitation.

The Fab or Fab', fragments of the anti FcyRII MAb were used for

immunoprecipitation studies from cell lysates of surface labelled K562 cells or Daudi cells (Figure 11A,B). Antibodies IV.3, CIKM5, 8.2 and 8.26 precipitated protein from K562 cells which migrated as a broad band on an SDS-PAGE gel with a Mr or 40-43kd. No material was precipitated with 8.7 antibody. A similar 40-43kD was immunoprecipitated with IV.3, 8.2 and 8.26 MAb from Daudi cells. In addition, aggregated human IgG and a rabbit polyclonal anti-FcγRII antibody immuno precipitated material of identical Mr to that of the FcγRII precipitated by the monoclonal antibodies (Fig. 11A, B).

#### Assay for circulating soluble FcyRII.

Using a two antibody sandwich radioimmunoassay normal individuals and a series of patients with autoimmune disorders (Systemic Lupus Erythematosis [SLE], Rheumatoid arthritis and Sjogren Syndrome) were tested for serum levels of soluble FcyRII. The results of this assay indicate that soluble FcyRII may be detected in normal individuals and patients with autoimmune diseases have significantly higher circulating levels of soluble FcyRII. The present of higher levels of circulating FcyRII is most evident in serum from individuals with Sjogren's Syndrome. Elevated levels were also detected in the circulation of patients with rheumatoid arthritis or Systemic Lupus erythematosis.

The production and characterization of four monoclonal antibodies to human FcyRII is useful in the study of cellular and circulating FcyRII and its role or presence in disease. The four new monoclonal antibodies (8.2, 8.26, 8.7, 7.30) clearly detect human FcyRII. Since they:-

- Bind specifically to mouse L cells transfected with human FcγRII cDNA but do not bind to the untransfected cells.
- 2) Show a tissue distribution that is identical to that expected for FcyRII.
- 3) Immunoprecipitate a 40 43 kd molecule that is the expected size of FcγRII and is the subject of Australian Patent Serial Number 595623.
- All antibodies partially or completely inhibit the binding of immune complexes (IgG- EA) to FcyRII positive target cells.

Clearly these antibodies detect FcyRII molecules. However it is clear that the antibodies fall into two discrete groups which when taken together with experiments

using two previously defined monoclonal FcyRII antibodies, indicate that at least four epitopes exist on FcyRII. The groups defined thus:-

- (i) 8.2, 8.26
- (ii) 8.7, 7.30
- (iii) IV.3
- (iv) CIKM5

These antibody groups can be distinguished from each other on the basis of (i) tissue distribution (ii) differences in the capacity to inhibit the binding of immune complexes, EA rosettes – (CIKM5 can only weakly inhibits EA rosetting whereas all other antibodies completely or significantly inhibit rosetting) and (iii) competitive inhibition assays indicate that the antibodies in the groups shown above detect four distinct epitopes. These distinct epitopes may be generated on mature cell surface FcγRII by post translation modification, such that the epitopes are carbohydrate determined; alternatively the antibodies detect protein epitopes in the peptide core.

It is clear that multiple forms of FcyRII have been defined (4-6, 8). However, these different forms have highly homologous extra cellular regions, wherein the amino acid identity was greater than 90% and the nucleic and amino acid sequence of the EC domains of the FcyRII, b1 and b3 are in fact identical. This high conservation of amino acid identify also results in high conservation of FcyRII function, in that all Fc receptors identified have the capacity to bind IgG immune complexes. To determine if the monoclonal antibodies described herein (8.2, 8.26, 8.7, 7.30) can distinguish between extracellular sequences of these multiple isoforms, transfection experiments were performed wherein cells were transfected with human FcyRII a clone 3.0 or transfected with the human FcyRII b1 clone (8). All monoclonal antibodies, 8.2, 8.26, 8.7, 7.30 bound to both FcyRIIa and FcyRII b1 and since the amino acid sequence of EC portion (both domains) of Fc RII b1, b2, b3 is identical (8) these MAb would detect FcyRIIb2 and b3 also.

Since the antibodies 8.26 and 8.7 detect distinct non overlapping epitopes, a capture tag radio immunoassay for the detection of soluble Fc receptor was developed. One monoclonal antibody was attached to the PVC plate and used to capture circulating Fc receptor in the blood of normal or diseased individuals. A

second antibody was used to detect the captured Fc receptor, this second antibody was tagged (eg. radio- labelled) on such a way that binding of this antibody could be detected. As shown in Figure 12 the second antibody was radiolabelled with I 125 and the specific binding of radiolabelled antibody to captured Fc receptor indicated the presence of circulating Fc receptor in blood derived from different individuals. In this study three groups of patients with autoimmune disease were studied: SLE, RA and Sjogrens syndrome. When compared to normal individuals all groups of patients with autoimmune disease has statistically significant high levels of soluble FcyRII. It is clear that this assay can be used to identify circulating soluble receptor and the assay can be adapted to be performed with other monoclonal antibodies or polyclonal receptor antibody. The second antibody can also be radiolabelled or conjugated to a flurochrome or enzyme and be used in photometric or colormetric assays, e.g. ELISA assays.

The detection of circulating FcyR may be of diagnostic use for patients with autoimmune or other diseases where high levels of receptor are of diagnostic or prognostic value or as an indicator of disease activity. Indeed, the patients afflicted by Sjorgens Syndrome and the diseases indicated (SLE and RA) (Figure 12) have much higher levels of circulating receptor than normal individuals. Although the source of the circulating FcyRII is unknown it is clear that it can be detected in the circulation;. Furthermore, as all the MAb – detect all isoforms of FcyRII – i.e. FcyRIIa, FcyRIIb1 (and FcyRIIb2, b3) then the assay would also detect the circulating forms of these. However, the fact that the precise nature of the circulating FcyRII is unknown and does not compromise the use and application of the assay for the detection of soluble FcyRII.

The application of this assay for the detection of soluble Fc receptor is not restricted to use in assays of patient serum but is equally applicable for assays using other bodily fluids, eg. urine, saliva, synovial fluid, or faeces.

#### **Concluding Remarks**

We have demonstrated the principle that chimeric Fc receptors derived in such a fashion that they contain amino acid sequence from multiple Fc receptors can also retain the immunoglobulin binding functions of these receptors types. Also the

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identification of the function or active region of these receptors (ie. immunoglobulin interactive) regions also identifies a useful region of these molecules. Monoclonal antibodies were used to demonstrate the preservation of epitopes in these chimaeric molecules and detect soluble Fc receptor in the circulation of diseased individuals.

#### MATERIALS AND METHODS

#### Molecular genetic techniques.

Standard molecular genetic techniques were performed as described in references 13 and 14 and included; restriction digestion, electrophoresis and purification of DNA fragments, ligation, bacterial transformation, di-deoxy nucleotide sequencing, plasmid DNA preparation, nucleic acid phosphorylation and de-phosphorylation, hybridisation, Southern blots.

## Splice overlap extension (SOE).

Chimaeric cDNA (which encode chimeric Fc receptors) were produced by splice overlap extension (SOE) using three polmerase chain reaction (PCR) steps essentially as described (24,25). The first PCR was performed to amplify sequence that forms 5' section of the chimaeric cDNA. The second PCR amplified sequence that forms the 3' section of the chimeric cDNA. Oligonuleotides (500ng) hybridising to the 5' or 3' ends of the region to be amplified were mixed with the appropriate template (100ng) and 25 cycles were performed using a thermal cycler (see below). The sequence of oligonucleotides used in the PCR reactions were designed such that the oligonucleotide primer hybridising to the 3' end of the first PCR product (this makes up the 5' sequences of the chimeric cDNA) overlaps sequence contained within the oligonucleotide primer hybridising to the 5' end of the second PCR product (this makes up the 3' sequences of the chimaeric cDNA) (Figures 6,7 Table 3,4,9). Thus the two independently derived PCR products overlap at their 3' (PCR product 1) and 5' (PCR product 2) ends.

The splice overlap extension reaction (ie third PCR) generates the chimeric cDNA for subsequent manipulation and was performed using purified PCR products 1 and 2. Approximately 10ng of each of the PCR product was mixed with two oligonucleotide primers (used in PCR 1 and PCR 2) that hybridise to the 5' end of PCR product 1 or the 3' end of PCR product 2. Also the 5' oligonucleotides (NRI or

EG6) used in the third PCR contained a EcoRI recognition sequence for subsequent subcloning into the pKC3 vector. Similarly oligonucleotide EG5 that hybridises to the 3' end of the chimeric cDNA contained a Sall recognition site for subsequent subcloning of the chimaeric cDNA into the pKC3 vector. The PCR reaction were conveniently performed under the standard conditions i.e. oligonucleotides and template (quantities as required – see above) were mixed with 2.5U of Taq DNA polymerase in 10mM Tris-HCI pH 8.3, 50mMKCI, 1.5MgCI<sub>2</sub> (Varied according to oligonucleotide primer combination). Twenty-five cycles were performed each cycle consisting of denaturation at 94°C for 1 minute, annealing for 2 minutes at appropriate temperature and extension at 72°C for 3 minutes.

#### Cloning of hFceRI cDNA.

1st strand cDNA was produced from human PBL (26). PCR was performed on 1st strand cDNA using oligonucleotide primers MDH13 5TTAGATCTCAGCACAGTAAGCACC3' which hybridizes to nucleotides position 1 to 17 of FceRI (non-coding strand) (12) and MDH14-5' TTTAGATCTAAATTGAACATCTCTTTTAC 3' (positions 1042 TO 1062). Both oligonucleotides contain a Bgl-II sit at their 5' ends for subcloning of the PCR product into pKC3 vector. The nucleotide sequence of the cloned FcRI DNA was determined by dideoxy-nucleotide sequencing (13,14) and is shown in Figure 7. FcyR cDNA.

Mouse FcyRII and FcyRI cDNA used herein have been previously described (4, 5, 7). The human FcyRIIa cDNA sequence used herein has been described (6, Australian Patent Serial No. 595623) and the sequence shown in Figure 6.

Production of Monoclonal Antibodies. BALB/c mice, 8-10 week females, were immunised by weekly intraperitoneal injection of whole cells (0.5mls of 10<sup>8</sup>-10 cells/ml) for a least 3 weeks with either the K562 cell line or L-cells transfected with the human FcγRII 3.0 cDNA (Tf3.0) (6). Cell fusions using the NS-1 myeloma cell line were performed as described (29). Hybridoma supernatants were screened for MAb by two stage rosetting of target cells using sheep anti-mouse immunoglobulin coupled to SRBC via CrC12 (28). Target cells used for screening were both K562 and Tf3.0 cell line. Positive hybridmas were then cloned by limiting dilution at least

twice and then grown as ascites tumors in (CBAxBALB/c)F1 mice primed with pristane. MAbs were purified from ascites using Protein A Sepharose chromatography. Other MAb used in the study included IV.3 (IgG2b) and CKM5 (IgG1) both standard anti CD32 MAb (22, 23). Anti-Ly2,1 antibody 49.11.1 (IgG2a) and 49-17.1 (IgG), and 5084-4.1 (IgG2a) anti-Ly-12.1 (IGg2A) used as a negative control antibodies.

#### Quantitative immunofluorescence.

Binding of MAb to haemopoletic cell lines was quantitated on a Facscan by immunofluorescence. 25ul of ascites or serum (1:400 dilution) or aggregated human IgG in PBS was added to target cells (5x10<sup>6</sup> cells/ml) in PBS-BSA 0.5% and incubated for 45 minutes on ice. The monocyte cell lines U937 and HL-60 were pre-incubated with 25 ul of 5mg/ml of human immunoglobulin to block non specific Fc binding to high affinity FcγRI. Cells were washed and resuspended in fluorescein isothyiocanate (FITC) sheep anti-mouse Ig F(ab)'2 (silenus) 1:50 dilution) was then incubated on ice for 30 minutes in the dark. Cells were washed and resuspended in 0.5ml PBS-BSA 0.5%.

#### Inhibition of EA Rosetting.

The ability of the MAb to block Fc binding was determined by inhibition of rosetting with polyclonal rabbit antibody coated erythrocytes (EA). The inhibition assay was performed as follows: 50ul of the blocking antibody ascites or serum was serially diluted and incubated with 50ul of FcγR bearing target cells at a concentration of 5x106/ml for 45 minutes on ice. Cells were washed free of excess antibody. 50ul of EA was added to the cell and spin at 200g for 4 minutes. The EA's and cells were incubated for 30 minutes on ice. Cells were stained with Ethyl Violet and a typical field of 100 cells was assessed for rosette formation (at least 5 RBC bound or 50% of cell surface covered). Target cells used were K562, Daudi and Tf3.0. Sensitisation of sheep erythrocytes with antibody was performed as described (5, 29).

#### Competitive inhibition Assays.

Purified antibodies (100ug) were labelled with carrier free I<sup>125</sup> using the Chloramine T method 29) but labelling performed for 30 seconds on ice with a

Chloramine T concentration of 1mg/ml. Free iodine was removed on a Sephadex PD-10 column (Pharmacia). Flexible microtitre plates were coated with 5% skim milk overnight to reduce non specific binding. Serial two-fold dilutions starting at a final concentration of 25 ug/ml of unlabelled antibody (25 ul) was mixed with 25 ul of a fixed predetermined dilution of radiolabelled antibody. Finally 50 ul of target cells 10<sup>7</sup> cells/ml) was added to the mixture of labelled and unlabelled antibody, and incubated at 4°C for 4 hours. Cells were washed and assayed for bound radioactivity. Value of radioactivity bound to cells in absence of unlabelled antibody was taken as 100% binding. The % inhibition was calculated as follows:-

 $(max. cpm) - (xcpm) \times 100$ 

max.cpm 1

where x=cpm of assayed cells and max.cpm = 100% binding value. Target cells used where FcyRII cDNA transfected L-cells and K562 cell.

Unlabelled irrelevant antibodies 49-11.1 or 49.17.1 antibodies were used to assess non-specific binding.

## Immunoprecipitation.

Cell suspensions (5x10<sup>7</sup> to 10<sup>8</sup>) were surface labelled with carrier free I<sup>125</sup> using a modified lactoperoxidase method (29). I<sup>125</sup> (10ul of 1mCi/ml; Amersham) and lactoperoxidase (80ul, 1mg/ml PBS; Sigma) was added to the cell suspension and surface radioiodination initiated by sequential addition of H<sub>2</sub>O<sub>2</sub> (BDH) (20ul) of 1:27,000, 1:900, 1:2000, 1:1000 dilution of 30% v/v solution in PBS) at 2 minute intervals. The reaction was stopped by addition of cold PBS and the cells washed twice in PBS. Cells were then solubilised in lysis buffer (0.01M Tris.Ha),.15M NaCI, 0.5% NP 40, 1mM EDTA and 1mM Phenyl sulphonylfuoride pH=8) at 4°C for 2 hours. Nuclei were removed by centrifugation at 10,000g for 10 min. Free iodine was removed on a Sephadex PD-10 column. Immunoprecipitaion was carried out by incubating 50ul antibody (Fab/Fab'2) conjugated sepharose beads (washed in lysis buffer) with 1ml of cell lysate at 4°C overnight. Polyclonal antibody or HAGG were incubated with 50ul of protein A conjugated Sepharose and then with 1ml of cell lysate at 4°C. The following day beads were washed in lysis mix, dissolved in SDS-PAGE sample buffer and analysed on a 10% SDS-PAGE gel under reducing

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conditions (29). Dried gels were autoradiographed.

Assay for Human Soluble FcyRII Using a Two Antibody Sandwich Radioimmunoassay.

Two monocolonal antibodies (8.2 and 8.26) to human Fc RII detecting different epitopes were purified from ascities using Protein A Sepharose chromatography (29). Flexible 96 well PVC microtitre plates (Costa) are coated with the first antibody 8.26 using 50–70ul per well at a concentration of 10ug/ml of antibody diluted in 0.005M carbonate/bicarbonate coating buffer pH=9.6 and incubated for 2 hours at 37°C. The plates are then washed three times by flooding the plates in PBS/0.05% Tween 20. The remaining sites in the wells are then blocked by coating the wells with 200 ul of 5% BSA and incubating for 1 hour at 37°C to reduce non-specific binding. The plate is then washed three times in PBS/0.05% Tween 20. Serum samples are diluted 1:4 in 2% BSA and 50ul is added to each well and incubated at 4°C overnight. The plates are then washed three times in PBS/0.05% Tween 20. The second radiolabelled antibody (8.7) was then added. 50ul of <sup>1125</sup> labelled 8.7 (50x10<sup>4</sup> cpm/50ul) is added to each well and incubated for 4 hours at 4°C. The plates were washed four times by flooding in PBS/0.05% Tween 20 and individual wells were cut and counted in Gamma counter.

#### Transfections.

Transfection of plasmid DNA into COS-7 cells was performed using DEAE dextran as described (10, 13, 14).

#### Monoclonal antibodies.

The monoclonal anti-FcyRII antibody IV.3 was obtained from Dr Clark Anderson (Ohio State university) and the CIKM5 antibody from Mr Glen Pilkington (Cancer Institute, Melbourne).

The IgG anti-TNP antibodies used have previously been detailed (5). The IgE anti-TNP antibody, TIB-142 was obtained from the ATCC (Maryland, USA). The monoclonal anti-glycophorin antibody was obtained from Dr Leonie Ashman (University of Adelaide).

The 2.4G2 antibody has been described (30).

Fab'<sub>2</sub> or Fab fragments of antibodies were produced as described (29).

#### Polyclonal anti-FcyRII antibodies.

Rabbit antibody detecting human FcyRII was produced by immunisation of New Zealand White rabbits with purified FcyRII fusion protein. The rabbit were immunized at regular intervals five times 3-5mg of fusion protein. The first immunisation was performed intradermally in multiple sites in complete Freunds adjuvant and subsequent immunisations in incomplete Freunds adjuvant.

The bacterial fusion protein was produced using the pATH21 vector (31) by cloning the large Pst-1 - XbaI fragment of FcyRIIa CDNA (Figure 6) into the Pst1 site and XbaI sites of pATH21. The XbaI site being introduced by PCR mutagenesis into the FcyRLLa cDNA at position 627-632.

The induction and purification of the fusion protein by electrophoresis was performed as described (31)

#### Scatchard Analysis.

Scatchard analysis was performed as described (32) using monomeric mouse I<sup>125</sup> - lgG2a prepared by incubating 100ug of lgG2a (in 50ul of 0.3M phosphate buffered saline pH7.4) with 500uCi Na<sup>125</sup> (Amersham UK) in the presence of chloramine - T Merck) at 1mg/ml in PBS pH7.4, for 30 sec. on ice. Labelling reactions were stopped with 50ul of 2.4mg/ml sodium metabisulphite and 10mg/ml tyrosine; free 125 was removed from labelled protein by passage through a PD-10 Sephadex G-25 column (Pharmacia, Uppsala, Sweden. COS cells transiently transfected with FcR cDNAs were harvested 48hr after transfection, washed twice in PBS containing 0.5% BSA and resuspended to 2x10<sup>6</sup>/ml in L15-0.05% BSA for use in Scatchard analysis. 50ul aliquots of cells were incubated with 50ul serial dilutions of <sup>1125</sup> - IgG2a in L15 medium for 60 min at 25°C with periodic aggitation. Cells were then pelleted by centrifugation through a 3:2 (v/v) mixture of dibutylpthalate and dioctylphalate oils (Fluka Chemika, Switzerland) and cell bound <sup>125</sup> –lgG2a assayed. Parallel experiments were performed under identical conditions in the presence of 100 fold excess of unlabelled lgG2a to correct for non-specific lg binding.

#### <u>Oligonucleotides</u>

Oligonucleotides were synthesised as described (27) end are listed in Table 9.

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The invention will be further described with reference to the following figures:

Figure 1

Mutagenesis of FcγRI. PCR was employed to introduce an Apa I site between D2 and D3 of FcγRI cDNA to facilitate domain exchange with FcγRII cDNA. Two PCR reactions were performed, one to produce a fragment containing D1 and D2 coding regions and a second PCR to produce a fragment containing D3, the transmembrane and cytoplasmic tail coding regions. Each of the fragments were engineered to have an Apa I site introduced into the FcγRI sequence between the D2 and D3 coding regions – through the use of overlapping and partly complementary oligonucleotide PCR primers (the Apa I site is indicated by solid circles). Each PCR reaction also employed an oligonucleotide primer containing a Sal I site (solid box), thus producing FcγRI DNA fragments containing Sal I and Apa I sticky ends suitable for construction of the chimeric receptor cDNAs (see Fig. 2).

PCR conditions: The complementary oligonucleotide designed to introduce the Apa I site had the following sequences:

MDH1 5' GGTGAA GGGCCC TTTCACCGTGATGG 3'

MDH2 GGTGAAAGGGCCC TTCACCACGCCAG 3'

MDH1 corresponds to nucleotides 573 to 599 and MDH2 corresponds to nucleotide 580 to 605 of the FcyRI cDNA sequence.

Boxed sequence contains the Apa I site.

The oligonucleotides containing the Sal I site were as follows:

MDH3 5' TTT GTCGAC ATGATTCTTACCAGCCTTTGGAGATG 3'

MDH4 5' TTT GTCGAC CCCCGGGGATCCTCTAGAGTCGAC 3'

The boxed sequence contains the Sal I site. MDH3 hybridizes to FcγRI 5' untranslated sequence from position 1-26 the first 8 bases of MDH3 contain 3 spacer nucleotides and 5 bases of the Sal I sequence.

MDH4 hybridizes to the pGEXII vector sequence flanking the 3' end of FcyRI cDNA insert.

The two PCR employed the oligonucleotide pairs MDH1 and MDH3 to produce the D1 and D2 fragment and MDH2 with MDH4 to produce the D3, tm and cytoplasmic tail coding fragment. PCRs were performed under the following conditions: lng of FcyRI cDNA (cloned in the vector pGEXII) was used as a template for amplification of the two fragments outlined above. 25 cycles were performed using a Perkin-Elmer-Cetus DNA thermal cycle in the presence of 500ng of each oligonucleotide primer, 10mM Tris-C1 ph 8.3, 50mM KC1, 1.5mM mgCl<sub>2</sub> and 2.5 units of taq polymerase (Amplitaq – Perkin – Elmer – Cetus). The PCR products were extracted with phenol: chloroform and ethanol precipated. The precipates were dried, dissolved in buffer and digested with Sal I and Apa I, extracted, precipitated and used in subsequent cloning steps see Fig. 2.

# Figure 2

Construction of expressable chimeric FcR cDNA's.

Chimeric cDNA's were generated by replacing FcyRII coding domains in the expression vectors with FcyRI coding domains generated by PCR. (A) FcyRII cDNA was subcloned into the Pst I site of the expression vectors pKC3 and pKC4 (derived from the parental vector pKO-neo Van Doren, et al J. Virol. 50 (1984) in expressable orientation [pKC3 and pKC4 differ only in the orientation of the polylinker polycloning sites such that the Sal I site flanks the 5' or 3' end of the cDNA insert in pKC3 of pKC4 respectively].

- (B) To produce the FcγRI-II chimera (consisting of D1 and D2 of FcγRI linked to the transmembrane and cytoplasmic coding regions of FcγRII) D1 and D2 of FcγRII were removed from pKC3-FcγRII by digestion with Sal I and Apa I and (C) replaced with a Sal I Apa I fragment containing D1 and D2 of FcγRI derived by PCR (see figure 1).
- (D) Similarly, for generation of the FcγRII-I chimera (consisting of d1 and d2 of FcγRII linked to D3, the transmembrane and cytoplasmic coding domains of FcγRI) the cDNA sequences encoding FcγRII transmembrane and cytoplasmic regions were removed by digestion with Sal I and Apa I, and (E) replaced with Sal I Apa I fragment containing D3, the transmembrane and cytoplasmic coding domains of FcγRI derived by PCR (see Figure 1).

# Figure 3

Nucleotide and amino acid sequence of chimeric Fc receptors. Nucleotide positions are numbered in decades below the line in a 5' to 3' direction. Untranslated

sequence is shown in closed type.

- A. Chimeric FcyRI/II. Fc RI derived cDNA sequence from positions 9 to 594, FcyRII from 595 to 1244.
- B. Chimeric FcyRII/I. FcyRII derived cDNA sequence from positions 1 to 662, FcyRI from 663 to 1348 between FcyRI and FcyRII domains. The Apa I restriction, site GGGCCC in both chimeric sequences is underlined. Sal I sites introduced into FcyRI sequence by PCR to facilitate cloning are boxed.

# Figure 4

IgG immune complex binding to chimaeric FcYR using antibody sensitized erythrocytes (EA). COS-7 cells were transfected with cDNA encoding FcYRI (A,F,G); FcYRII (D); (FcYRI/II (C,I,J); FcYRII/I (B); vector only (E,H). The transfected COS cell monolayers were tested for Fc binding using erythrocytes snesitised with antibody. Rabbit IgG-EA were used in (A-E) and mouse IgG1-EA (G,H,J) and mouse IgG2a-EA (F,I). Methods described in Materials and Methods Section.

# Figure 5

Scatchard analysis of IgG2a binding to FcyRI transfected COS-7 cells. Data has been subjected to linear regression analysis.

# Figure 6

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Nucleotide and deduced amino acid sequence of human FcyRIIa (6). Amino acid positions are numbered above the line in decades commencing at the proposed amino acid terminal end. (Signal sequence is numbered from residue -34 to -1). Nucleotide positions are numbered at the end of the line. Oligonucleotide primers used in the PCR reactions for construction of the chimaeric cDNA are positioned at their priming sites with 5' to 3' direction indicated by half arrow heads. Solid lines represent oligonucleotidessequence derived from the FcyRII cDNA template, dotted lines represent oligonucleotide sequence derived from FceRI cDNA template (Figure 7). Solid boxes represent 5' terminal EcoRI sites, solid circles 5' terminal Sall sites. Figure 7

# Nucleotide and deduced amino acid sequence of human FceRI. Amino acid positions are numbered above the line in decades commencing at the proposed amino

acid terminal end (signal sequence is numbered from residue -25 to -1). Nucleotide positions are numbered at the end of the line. Oligonucleotide primers used in the PCR reactions for constructions of the chimaeric cDNA are positioned at their priming site with 5' to 3' direction indicated by half arrow heads. Dotted lines represent oligonucleotides sequence derived from the FcεRI cDNA template, solid line represents oligonucleotide sequence derived from FcγRII cDNA template (Figure 6). Solid boxes represent 5' terminal EcoRI sites.

#### Figure 8

Rosetting of COS-7 cells transfected with FcR cDNA. COS 7 cell monolayers were transfected with human FcyRII cDNA (A,B); Chimaera 1 cDNA (C,D); Chimaera 2(E,F); Chimaera 3 (G,H); chimaera 4 (I,J). Seventy-two hours later the binding of antibody sensitised erythrocytes was assessed using IgE-EA (A,C,E,G,I) or IgG-EA (B,D,F,H,J).

#### Figure 9

Fluorescence histograms of immunofluorescence of anti-FcyRII monoclonal antibody staining of Daudi cells of U937 cells as indicated. Horizontal axis represents fluorescence intensity (log scale) the vertical axis represents relative cell number.

#### Figure 10

Competitive binding of unlabelled monoclonal anti-Fc RII antibodies. Competitive assay performed using L-cells transfected with human FcyRII cDNA HFc 3.0 Radiolabelled antibodies are (A), IV.3; (B), CIKM5; (C), 8.2; (D), 7.30. The cold competitors are shown, Ly-2.1 (49-11.1); IV.3; CIKM5; 8.2; 7.30.

# Figure 11

Immunoprecipitation of FcyRII from surface labelled K562,(A) or Daudi,(B) cells. Molecular weight standards (Mrx10<sub>3</sub> (Std) are indicated on the left side of the figure. Antibodies (Fab or Fab'2 fragments) used are IV.3, CIKM5, 8.2, 8.7, 8.26. Human aggregated IgG (HAGG) intact Ly-12.1; 1705(5084-4.1). The negative control for the polyclonal antibody was immunoprecipitation from FcyRII-CEM cells.

# Figure 12

Detection of circulating FcYRII in the serum of normal or diseased individuals. (A) mean cpm ± 1 S.E. of bound anti-Fc RII in the patient groups indicated. P values are calculated for comparison of patient groups and normals. The numbers (n) of individuals tested within each patient group are indicated. (B) Dot plot of data obtained for individual patients within each group. Each point represents number of cpm (labelled anti-Fc RII antibody) bound.

The entire contents of the provisional specification lodged with Australian Patent Application of which this is the complete specification is hereby imported into this specification and forms part of the disclosure of this specification. The claims form part of the disclosure of this specification.

#### **DEFINITIONS:**

### Extra functional region:-

Part of a molecule, having function other than immunoglobulin binding. Functional region:—

Part of molecule capable of interacting with immunoglobulins or their fragments whether the immunoglobulins or fragments are monomeric in nature, aggregated or immune complexes.

### Immunoglobulin binding molecule:-

Any molecule capable of interacting with immunoglobulins or their fragments whether the immunoglobulins or fragments are monomeric in nature, aggregated or immune complexes.

#### Functional equivalents thereof:-

Variants on the materials the subject of this specification are possible:-

- A. Sequence Variation
  - The nucleotide sequences encoding the receptor can be variable:-
- 1. Because of the degeneracy of the genetic code nucleotide change does not necessarily bring about a change in the amino acid encoded.
- 2. Two or three nucleotide changes can give rise to the same amino acid.
- 3. Changing one or two nucleotides may result in a conservative amino acid change unlikely to greatly affect the function of the protein.
- 4. Allelic variations. Variations in nucleotide sequence and resultant amino acid sequences of the encoded protein may occur between individual members of the same species. These variations arise from changes in the nucleotide sequences encoding the protein. Thus different forms of the same gene (called alleles) give rise to protein of slightly different amino acid sequence but still have the same function.
- 5. Proteins having the same function, e.g. immunoglobulin binding, may arise from related genes. Many protein gene families have been described.
- 6. Variation may be intentionally introduced by:-
- (a) Mutating cloned cDNA or genomic DNA by point mutation, rearrangment or insertion of related or unrelated DNA into the cDNA or genomic clones

- encoding the functional protein. Such mutated (variant) clones can be used to generate variant proteins or peptides.
- (b) By enzymatic cleavage of the protein (from either in vitro synthesis or normal cell synthesised protein) with or without repair/rearrangement of the cleavage products.
- (c) By chemical modification.
- (d) By irradiation.

# Greek symbols:-

- a alpha
- γ gamma
- δ delta
- ε epsilon
- μ mu .

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Binding of IgG subclasses to chimeric receptors

TABLE 1

Transfecting	g DNA	Bine	ding of	IgG*		
			. 1	Mouse Ig	Gsubclas	ses <sup>t</sup>
	Human	Rabbit	G1	G2a	G2b	G3
	IgG**	IgG <sup>t</sup>				*.
FcRI	+	+	_	+	_	-
FcRII	+	+	+	+,	+	+
FcRI-II	+	+	+	+	+	+
FcRII-I	+	+	+	+	+	+
NIL	_	<b>-</b> ·	-	-	-	-

- \* Ig binding assessed by EA rosetting for rabbit and mouse lgG. And
- Ig binding level where >10 IgG coated erythrocytes bound per adherent Cos cell.
- \*\* binding assessed by immunofluorescence using heat aggregated-human poly clonal IgG and quantified by FACS analysis.

Erythrocyte - antibody complexes were prepared as follows:

- (i) Rabbit IgG sensitised erythrocytes were prepared as in Figure 1.
- (ii) EA coated with specific mouse IgG subclasses were produced by using anti-TNP isotype specific monoclonal antibodies (of the IgG1, IgG2a, IgG2b subclasses). Sheep red blood cells were sensitised with TNP by incubating a 10% SRBC suspension with 7 volumes of TNP in PBS (Phospate buffered saline) for 20 minutes at room

temperature. Sensitised cells were then washed twice in PBS+0.5% BSA and resuspended to - a 2% suspension upon which an equal volume of antibody was added and incubated for 60 minutes at room temperature.

(iii) Human red blood cells directly sensitised with IgG1 of IgG3 anti-glycophorin monocional antibodies as described above. EA's were used in rosetting assay again as outlined in Figure 4 legend.

Assays were performed directly in culture dishes on COS - 7 cells transiently transfected with FcR expression constructs (Fc RI in the pGEXII vector (Ref), Fc RII in pKC3, Fc RII-I in pKC4, Fc RII in pKC3), as described EA's were then washed twice in PBS-0.5% BSA again resuspended to a 2% suspension for use in rosetting following an approach outlined in Figure 3 legend.

Immunofluorescence was performed on transiently transfected cells. Polyclonal human IgG (5 mg/ml) was aggregated at 65°C for 30 minutes. Binding and immunofluorescence studies were performed as described (29).

Detection 2.4G2 epitope on chimeric receptors by blocking of EA rosetting

TABLE 2

Transfecting DNA	% Blocking EA rosetting*
FcRI	
FCRII	80
FcRI-II	80
FcRII-I	80
Vector only **	0

- \* % blocking relative to transfected cells not treated with 2.4G2 antibody.
- \*\* Cos cells were transfected with pKC4 vector DNA.

Blocking assay were performed on Cos 7 cells transiently transfected with FcR expression constructs (see Table 1). Transfected cells in culture dishes were incubated whilst adhered with Fab fragments of the anti-Fc RII monocional antibody 2.4G2 (Unkeless; J.C. Exp. Med 150: 580-596, 1979) for 60' at 4°C. Cells (in dishes) were then washed x2 with L-15-0.5% BSA medium, and rabbit EA added to dishes. Rosette formation was assessed as outlined to Figure 3 legend.

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CDNA IN THIS STUDY	ORIGIN OF AMINO ACID RESIDUES*,†			$[\epsilon(-25 \rightarrow 169), \gamma (170 \rightarrow 281)]$	$[\gamma(-34 \rightarrow 86), \epsilon (87 \rightarrow 169), \gamma(170 \rightarrow 281)]$	$[\epsilon(-25 \to 86), \gamma (87 \to 281)]$	$[\epsilon(-25 \rightarrow 128), \gamma (129 \rightarrow 281)]$	$[\epsilon(-25 \rightarrow 86), \gamma (87 \rightarrow 128), \epsilon(129 \rightarrow 169), \gamma(170.281)]$	$[\gamma(-34 \rightarrow 86), \ \epsilon(87 \rightarrow 128), \ \gamma(129 \rightarrow 2)$	$[\gamma(34 \rightarrow 128), \epsilon (129 \rightarrow 169), \gamma (170 \rightarrow 281)]$	D1 EC domain 1: D2 EC domain 9: TM transmembrane: CT evtonlasmic tail is derived from FosBl: v. derived from EcvB
FCR CDNA AND CHIMAERIC FCR CDNA IN THIS STUDY	BECEPTOR COMPOSITION", ** D1 D2 TM/CT									WINN WIN	o. TM transmembrane: CT cytoplasmic
TABLE 3	ORIGIN OF DOMAIN* D1 D2 TM/CT	ພ ພ	, , , ,	ک ع	γ γ	٠ ٠ ٠	3 ع	λ 3/k 3	γ ε/γ γ	γ γ/ε γ	oin 1. Do EC domain
	CHIMAEBA NAME	FceRI	FcyRII	-	2	· •	4	ĸ	9	7	1 FC - 1

FcR was determined by dideoxy nucleotide sequence analysis (12, 13). Amino acid residues of the receptors. Numbering indicates the amino acids derived from FcεRI (ε) or FcγRII (γ) and numbers correspond to amino acid residues in Figures 6 and 7. derived from FcyRII. Origin of leader sequences not shown. The nucleotide and amino acid sequence of all native and chimaeric D1, EC domain 1; D2, EC domain 2; TM, transmembrane; CT, cytoplasmic tail, ε, derived from FcεRI; γ, derived from FcγRII. Diagramatic representation of origin of the regions of the Fc receptors. Unshaded regions derived from FccRI; shaded regions

TABLE 4 CONSTRUCTION OF CHIMAERIC CDNAS

CHIMAERA NAME	ORIGI	ORIGIN OF DOMAINS	MAINS	OLIGONUCLEOTIDE Pair I (Template)*	Oligonucleotide Pair II (Template)*
	01	D2	TM/CT		
FcyRII	~	. ~	. ~		
FceRi	ယ	ພ	ω		,
-	ພ	ພ	٦,	EG6 + EG1 (ForRI)	EG2 + EG5 (FcyRII)
2	~	ພ	-ح	<u>NR1</u> + EG11 (FcyRII)	EG10 + <u>EG5</u> (Chimaera 1)
က	ယ	~	٨	EG6 + EG9 (FccRI)	EG8 + EG5 (FcyRII)
4	ω	٤/٦	٨	<u>EG6</u> + EG14 (Fc:RI)	EG15 + <u>EG5</u> (FcyRII)
n L	ພ	3/£	٨	EG6 + EG12 (Chimaera 3)	EG13 + <u>EG5</u> (Chimaera 1)
9	۲	٤/٧	٨	NR1 + EG14 (Chimaera 2)	EG15 + EG5 (FCyRII)
7	٨	3/κ	۲	<u>NR1</u> + EG12 (FCyRII)	EG13 + EG5 (Chimaera 1)

Underlined oligonucleotides used in SOE reaction; templates for PCR reactions shown in parentheses.

ANTI FCYRII MOAD BINDING!

PROPERTIES OF NATIVE FCYRII, FCERI AND CHIMAERIC FCR TABLE 5

IMMUNE COMPLEX BINDING"

CHIMAERANAME	OBIG	NOFD	ORIGIN OF DOMAIN*	ð	<u>ng</u>	8.7	7.30	8.2	8.26	17-3	CIKM5
	10	D2	TM/CT	•							
FceRI	ω	ω	ω.		1			•	•	ı	t
FcyRII	<b>,</b>	٨	<b>~</b>	+	•	+		+	. +	+	+
	ယ	မ	۸	•	+	•		•	,	•	
2	*	ພ	ယ	ı	+	1	1	. 1.	ı	ı	ı
3	ພ	ح	٠,	+		+		1	•	+	
4	ω	٤/ ٢	٨	+		9		2	Q	2	2
വ	<b>ພ</b> .	٤/٦	٠.	ı	•	2		2	2	2	9
9	٨	٤/ ٢	۸	+	+	Q		2	Q	2	2
7	٨	3/6	<b>~</b>	,		2		2	2	2	<del>Q</del>

D1, EC domain 1; D2, EC domain 2; TM, transmembrane; CT, cytoplasmic tail; ε, derived from FcεRI; γ, derived from FcγRII.

immune complex binding assessed by EA rosetting using IgG and IgE sensitised erythrocytes. + = formation of rosettes; - = no rosette formation.

Binding assessed Binding of monoclonal anti FcyR antibodies (MoAb) to cells transfected with native or chimaeric Fc receptors. Binding by immunofluorescence and visualisation by fluorescence microscopy. +, fluorecent staining; -, no staining; ND, not determined.

CHARACTERISATION AND TISSUE DISTRIBUTION OF MONOCLONAL ANTIBODIES TABLE 6

MoAb	Immunoden	la Class	SS					Target Cells	Cells						
			Tf3.0 L-0	L-cells	sells K562 DAUDI RAJI	DAUDI		EBV	1937	ThP1	HL-60	OEW EW	JURKATT	U937 ThP1 HL-60 CEM JURKATT MOLT4 PEER	出
								B cells							
17.3	K562	γ2 b	+++++		+++++	+		+	+++++	++++	+				Ï
<b>CIKM5</b>	K562	۲	++++	•	+ + +	+	+	+	+ + +	+	+				
8.2	K562	۲۸	+ + +		+ + +	+	•	+	+++	+ + +	+			•	
8.7	K562	ሃ	++++		+ + +	+ + +	+		+	+ + +	+	1		,	
8.26	K562	γ2 b	++++	•	++++	++	+	+ +	+ + +	+ + +	+				
7.30	Tf3.0	71	++++	•	++++	+ + +	+	+	++	+ + +	+	1			
														-	1

% reactivity with haemopoietic cell lines and human FcyRII transfected L cells compared to the negative control mAb 49.11.1. Ascites or serum was used at dilution of 1:400 0-10% reactivity = Negative (-); 10-50% reactivity = +; 50-75% reactivity = ++; 75-100% reactivity = +++. The control mAb 49.11.1 was negative in all cases.

Table 7

INHIBITION OF IMMUNE COMPLEX BINDING (EA-ROSETTING)

Target Cell mAb K562 Tf3.0 DAUDI 100 **IV.3** 100\* 100 CIKM5 44 0 0 8.2 100 100 100 8.7 78 88 100 8.26 100 100 100 7.30 44 75 100 49.11.1 0 0 0

Percentage of inhibition of EA rosette formation using all indicated antibodies at a 1:2000 final concentration of ascites or serum. Target cells indicated.

TABLE 8

COMPETITIVE BINDING ASSAY\*

		RADIOLABEL	LED ANTIE	BODY	·-	
Cold	IV.3	8.2	8.26	CIKM5	8.7	7.30
Competitor						
						<del></del>
IV.3	+	+	+	+	+	+
8.2	+	+	+	+	-	•
8.26	+	+	+	+	•	-
CIKM5	±	±	±	+	±	•
8.7	±	-	-	±	+	+
7.30	±	•	-	±	+	+

#### K562 is the target cell

- +; % inhibition equivalent to that obtained with the same unlabelled antibody e.g. unlabelled IV.3 inhibition of labelled IV.3.
- ±; partial or minimal inhibition of radiolabelled antibody
- -; no inhibition of radiolabelled antibody.

OLIGONUCLEOTIDES USED IN THE GENERATION OF CHIMAERIC FCR TABLE 9

NAME	SEQUENCE 5' - 3'	NUCLEOTIDE POSITION ON FŒRI/FCyRII cDNA** (5' to 3' of oligonucleotide)
EGI	5 GGCACTTGTACAGTAATGTTGAGGG3	FCYRII 628-621; FCERI 621-604.
253	5 CATTACTGTACAAGTGCCCAGCATG3	FceRI 612-621; FcyRII 621 to 635.
EGST	5 TTTGTCGACCACATGGCATAACG3	FcyRII 981-967
EG6†	5 TTTGAATTCAGCACAGTAAGCACC 3"	FccRI 6-22
EG8	5 TTCAGTGACTGGTGCTCCAG3"	FceRI 364-372; FcyRII 372-386
<u>EG9</u>	5 ACCAGCCAGTCACTGAAGACTTCC3	FcyRII 379-372; FceRI 372-357
EG10	5 CTTTCCGAATGGCTGCTCCTTCAG3	FcyRII 363-371; FccRI 373-387
<b>E</b> G11	5 AGCAGCCATTCGGAAAGCACAGTC3	FceRI 380-373; FcyRII 371-356
EG12	5 TACCAGTATTTCTGGGATTTTCCATTC3	FceRI 506-499; FcyRII 497-479
EG13	5 TCCCAGAAATACTGGTATGAGAACCAC3	FcyRII 489-497; FcERI 499-516
EG14	5 CGGGAGACTTGAGAGCTTCACCATC3	FcyRII 505-498; FccRI 498-481
EG15	5 CTCTCAAGTTCTCCCGTTTGGATCC3"	FceRI 491-498; FcyRII 498-514
NRIT	S TACGAATTCCTATGGAGACCCAAATGTCTC3	FcyRII 10-30

Underlined oligonucleotides complementary to coding strand.

Sequence numbers taken from Figures 6, 7.

#### CLAIMS:

- 1. A hybrid immunoglobulin binding molecule capable of binding to any one class or a plurality of classes of antibody molecules.
- 2. A hybrid immunoglobulin binding molecule according to claim 1 which is derived from any Fc receptor (FcR).
- 3. A hybrid Fc receptor molecule according to claim 2 comprising one or more functional regions of a first FcR linked to one or more functional regions of a second FcR or consecutive FcR.
- 4. A hybrid Fc receptor molecule comprising a polymer or a concatomer of heterogenous or homogenous functional regions.
- 5. A hybrid FcR according to any one of claims 1 to 4 wherein said functional regions are derived from different FcR molecules.
- 6. A hybrid FcR according to any one of claims 1 to 5 wherein said functional regions are derived from bacterial, mammalian or other origins.
- 7. A hybrid FcR according to any one of claims 1 to 6 wherein said functional regions are derived from human, mouse or a combination of human and mouse origins.
- 8. A hybrid FcR according to any one of claims 1 to 7 which is capable of binding any one of, or a plurality of antibodies selected from the classes IgG, IgM, IgA, IgD and IgE.
- 9. A hybrid FcR according to any one of claims 1 to 8 wherein said functional regions are derived from any one of FcR, FcγR, FcγR, FcαR, FcεR, FcμR or IgE binding proteins.
- 10. A hybrid FcR according to any one of claims 1 to 9 wherein the said functional regions are derived from FcyR or FceR being any one or a combination of the extracellular domains, or parts thereof, of FcyRI, FcyRII, FcyRIII, FceRI, FceRII or homologues or functional parts thereof.

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11. A hybrid FcR according to any one of claims 1 to 10 wherein said functional domains comprise one or more extracellular domains, or parts thereof, derived from FcyRII of mammalian origin in combination with any one extracellular domain derived from the following:-

mammalian FcγR
mammalian FcεRI,
mammalian FcεRII,
mammalian FcγRII,
mammalian FcγRIII,
mammalian FcαR,
mammalian FcuR, or
mammalian IgE binding proteins

- 12. A hybrid FcR according to any one of claims 1 to 11 wherein said functional regions comprise the extracellular domain or parts thereof derived from FcγRI being amino acids 1–171 linked to the transmembrane region and cytoplasmic tail of FcγRII being amino acids.
- 13. A hybrid FcR according to any one of claims 1 to 11 wherein said functional regions comprise extracellular domains or parts thereof, derived from FcyRII being amino acids 1-174 linked to the third extracellular domain, transmembrane region and cytoplasmic tail of FcyRI being amino acids 174-380.
- 14. A hybrid FcR according to any one of claims 1 to 11 wherein said functional region comprise an extracellular domain, or parts thereof, having greater than 90% homology to FcyRII or FceRI of human or mouse origin linked to an extracellular domain of FceRI of human or mouse origin.
- 15. A hybrid FcR according to any one of claims 1 to 11 wherein said functional regions comprise an extracellular domain, or parts thereof, having greater than 50% homology to FcyRII or FceRI of human or mouse origin linked to an extracellular domain of FceRI of human or mouse origin.
- 16. A hybrid FcR according to any one of claims 1 to 11 wherein said functional regions comprise an extracellular domain, or parts thereof, having greater than 35% homology to FcγRII or FcεRI of human or mouse origin linked to an extracellular domain of FcεRI of human or mouse origin.
- 17. A hybrid FcR according to any one of claims 1 to 11 wherin at least one functional region comprises an extracellular domain, or parts thereof, derived from FcyRII having the following amino acid sequence:—

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- Phe Ser Arg Leu Asp Pro Thr Phe Ser Tle Pro Gln Ala Asn His Ser His Ser Gly Asp Tyr His Cys Thr Gly Asn Ile Gly Tyr Thr Leu Phe Ser Ser Lys Pro Val Thr Ile Thr Val, or functional equivalent thereof.
- 18. A hybrid FcR according to any one of claims 1 to 11 wherein at least one functional region comprises an extracellular domain, or parts thereof, derived from FceRI having the following amino acid sequence:
  - Trp Leu Leu Gln Ala Ser Ala Glu Val Val Met Glu Gly Gln Pro Leu Phe Leu Arg Cys His Gly Trp Arg Asn Trp Asp Val Tyr Lys Val Ile Tyr Tyr Lys Asp Gly Glu Ala Leu Lys, or functional equivalent thereof.
- 19. A hybrid FcR according to any one of claims 1 to 18 wherein said molecule is not bound to a membrane and is soluble in physiological and or non physiological buffers.
- 20. A hybrid FcR according to any one of claims 1 to 18 wherein said molecule is linked to any transmembrane region and/or cytroplasmic tail.
- 21. A hybrid FcR according to claim 20 wherein said transmembrane region and cytoplasmic tail are derived from FcR.
- 22. A hybrid FcR according to any one of claims 1 to 121 being a chimera comprising a first extracellular domain of FcyRII, a second extracellular domain of FceRI and a transmembrane region and cytoplasmic tail of FcyRII.
- 23. A hybrid FcR according to any one of claims 1 to 21 being a chimera comprising a first extracellular domains of FcεRI containing amino acids 1 169 of FcεRI linked to amino acids 170-281 of FcγRII.
- 24. A hybrid FcR according to any one of claims 1 to 21 being a chimera comprising a first extracellular domain of FcεRI being amino acids 1-86 linked to a second extracellular of FcγRII being amino acids 87-169 linked to a transmembrane region and cytoplasmic tail of FcγRII being amino acids 170-281.
- 25. A hybrid FcR according to any one of claims 1 to 21 being a chimera comprising a first extracellular domain of FceRI containing amino acids 1-86 linked to part of the second extracellular domain of FceRI being amino acids 87-128 linked to part of the second extracellular domain of FcyRII being amino acids 129-169

linked to a transmembrane region and cytoplasmic tail of FcyRII being amino acids 170-281.

- 26. A hybrid FcR according to any one of claims 1 to 21 being a chimera comprising a first extracellular domain of FcεRI being amino acids 1-86 linked to part of the second extracellular domain of FcγRII being amino acids 87-128 linked to part of the second extracellular domain of FεRI being amino acids 129-169 linked to a transmembrane and cytoplasmic tail of FcγRII being amino acids 170-281.
- A hybrid FcR according to any one of claims 1 to 21 being a chimera comprising a first extracellular domain of FcγRII being amino acids 1-86 linked to part of the second extracellular domain of FcγRII being amino acids 87-128 linked to part of the second extracellular domain of FcεRI being amino acids 129-169 linked to the transmembrane and cytoplasmic tail of FcγRII being amino acids 170-281.
- 28. A hybrid FcR according to any one of claims 1-21 being a chimera comprising the first extracellular domain of FcyRII being amino acids 1-86 linked to the second extracellular domain of FceRI being amino acids 87-169 linked to a transmembrane and cytoplasmic tail of FcyRII being amino acids 170-281.
- 29. A hybrid FcR according to any one of claims 1-21 being a chimaera comprising the first extracellular domain of FcγRII being amino acids 1-86 linked to part of the second extracellular domain of FcεRI being amino acids 87-128 linked to part of the second extracellular domain of FcγRII being amino acids 129-169 linked to the transmembrane and cytoplasmic tail of FcγRII being amino acids 170-281.
- 30. A hybrid FcR according to any one of claims 1 to 29 further comprising one or more extra functional regions of FcR as hereinbefore defined.
- 31. The amino acid sequence comprising the hybrid FcR according to any one of claims 1 to 31.
- 32. A nucleotide sequence capable of encoding an amino acid sequence according to claim 31.
- 33. The nucleotide sequence of claim 32 comprising a cDNA or genomic DNA clone capable of encoding the amino acid sequence according to claim 31.
- 34. A method of producing hybrid immunoglobulin binding molecules

3

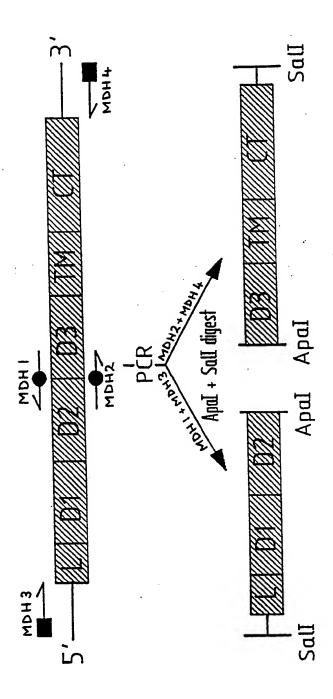
according to any one of claims 1 to 31 comprising the steps of linking different functional regions of different FcR molecules to construct a hybrid immunoglobulin binding molecule.

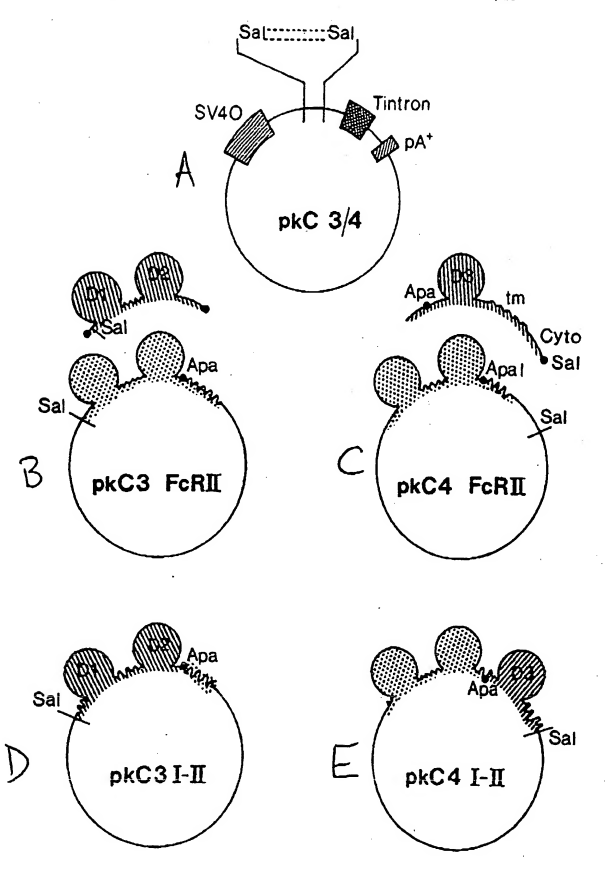
- 35. A method according to claim 34 comprising ligating a first FcR molecule at the transmembrane junction, ligating a second FcR molecule to obtain a suitable functional or non-functional region and linking said components.
- 36. A polyclonal antibody which reacts with any one of the hybrid FcR molecules as defined in claims 2 to 31, or parts thereof.
- 37. A monoclonal antibody which reacts with any one of the hybrid FcR molecules as defined in claims 2 to 31, or parts thereof.
- 38. A monoclonal antibody being of subclass IgG1 designated 8.2 as hereinbefore defined.
- 39. A monoclonal antibody being of subclass IgG1 designated 8.7 as hereinbefore defined.
- 40. A monoclonal antibody being of subclass IgG2b designated 8.26 as hereinbefore defined.
- 41. A monoclonal antibody being of subclass IgG1 designated 7.30 as hereinbefore defined.
- 42. Antibodies according to any one of claims 36 to 41 which detect a single epitope specific for FcR.
- A biological product comprising one or more hybrid FcRs according to any one of claims 2 to 31 or antibodies thereto according to any one of claims 36 to 41 for interaction with a hybrid FcR as defined in any one of claims 2 to 31.
- A biological product according to claim 43 wherein said interaction comprises any one or a combination of binding, detection, linking, inhibiting function.
- A biological product according to any one of claims 43 or 44 which is capable of being bound to fixed solid support being a tube, dipstick, multiwell plate or other form made from PVC or other material.
- 46. A biological product according to any one of claims 43 or 44 which is capable of being bound to biologically active materials including cytokines,

hormones, red blood cells, liposomes, dialysis membranes, etc.

- 47. A biological product according to any one of claims 43 or 46 which is capable of being labelled with any one of labelling enzymes, radioactive isotopes, metal or non-metal particles, fluorescent molecules, free radicals chemiluminescent or bioluminescent molecules.
- 48. A biological product according to any one of claim 43 to 47 further comprising anti FcR antibodies or fragments thereof.
- 49. A diagnostic test kit comprising the biological product of any one of claims 43 to 48.
- 50. An assay for the FcR being an immunoassay utilizing the antibody or biological product of any one of claims 36 to 48.
- 51. An assay according to claim 50 being a capture tag assay.
- A capture: tag assay according to claim 51 wherein the FcR is bound, and different epitopes of the said FcR are utilized for further binding and or antigenic functions.
- 53. A capture: tag assay according to claim 51 wherein the FcR is free.
- An assay according to any one of claims 50 to 53 comprising the steps of washing away unbound material, adding biological fluid (e.g. serum), incubating for ½-16 hours, washing unbound materials, adding a second labelled anti-human Ig (labelled for example with radioisotope; enzyme particle or fluorochrome), incubating for 1-24 hours, washing and determining the quantity of labelled antibody bound.
- A capture tag assay according to any one of claims 51 to 53 wherein said assay utilizes hybrid FcR as a standard or competitive inhibitor.
- 56. A method of regulating antibody production in vivo by binding a hybrid FcR according to any one of claims 2 to 30 to B cells.
- 57. The use of the method of claim 56 to control autoimmune disease.
- 58. The use of the method of claim 56 to control allergy reaction by affecting IgE control.
- A method of regulating allergic reactions by reacting hybrid FcR according to any one of claims 2 to 30 with IgE in vivo.

- 60. The use of the biological product of any one of claims 43 to 48 for plasmaphoresis to remove immune complexes or pathological antibodies.
- 61. The use of the biological product of any one of claims 43 to 48 in conjunction with biosensors.





# FIGURE THREE A

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		•						060	440	H	υH	<b>F</b> ==
	His CAC 180								S GCT 900	TCTCCCT 990	GGCCAGGATCTAGTGTCTCGTGGTCCAAGGGATGCTGTAGATATTAAAGAAACATCCAGAGTCACTTCTGTGAGTCCTGAAACCAACAACACAC 1021 1031 1041 1061 1061 1061 1071 1081 1081 1111	123 123
Val GTT 90	CCT	Ser		C C	CAC	TTG			Glu GAG		<b>S</b> CO	TACT
Val	61y 666	Ala GCC				Val GTA		SOL		TAG	NACC.	221 221
Glu GAA	Glu GAG 170	G1u GAG 60	7rp 766 350	AAT 440	11e ATC 530	Pro CCA 520	Gln CAG 710	61y 66C 800	Pro Lys CCC AAA 890	His Ile CAC ATT 980	TGA	LYCL
G1y GGG 80	Cys TGT	Pro Glu CCA GAG 260	Asp	Tyr Tac	ດ ຄຸດ	Leu Pro TTA CCA 620	Lys Ag	669 8		CAC	GTCC	LATE
Gly GGT	Trp	IIIe	Ash	Val GTG	Ser	Ser	Lys	Ser	Ala	Aan	1091	121
Val GTC	r Leu	Ser Agc	H1s CAC 0	3 Leu A CTG 430	Ser His AGT CAC 520	Arg Agg 10	Leu Lys CTC AAG 700	Gln Pro CAG CCC 790	GAA BO	GIn CAA 70	TTC	CCAC
Fro CCA	Thr 1	Tyr Ser Tat AGC 250	Ile ATC 34	Eys AAA 43	Ser AGT 52	Ser AGC 61	CTC	Gln CAG	GAA B	Tyr TAC 97	TCAC 81	ACTG 01
Va.	Val	Ser	GIn CAA	ABA	Leu CTG	SOL SOL	Tyr Tat	Arg Aga	CTO	ASP	AGAG 10	CCAC 12
Trp TGG	Aan	Pro	TTG	Lys	AAC	Lys	Val	TAC	Asp Gat	His	ATCC	CGAT
CTT	Glu / GAA / 150	Thr 1 ACG 240	Gln CAG 330	117 166 120	Thr ACC 510	200 000 000 000 000	110 110 690	Glu GAG 780	Pro CCT 870	GAG 960	AAAC 1071	CTAT 1191
Leu CTA 60	Lys AAG	Sor	Val	Gly GGA	Lye	959 969	Ser TCC	Gly GGT	Pro	ACA	AAGA	TCTG
CTG	Oln I	ATC	Pro	His	Lec	Ly.	Val	Val GTA	Aen Aat	GAA	ATTA 61	CTCT
Thr	Phe C	Gln CAG 230	Asp GAC 320	Cys TGT 410	11e ATT 500	Val GTG	CTA 680	GAA 770	TAC B60	Asp Glu GAT GAA 950	AGAT 10	AAGA 11
Thr ACA 50	ATT 1	Val GTT	Ser AGT	Arg Agg	Ala GCC 5	Thr Acc 5	ATC	G1u GAG	Pro CC.	Asp GAT	crer	ACTO
Leu	Ser	Ala	Pro	TTG	Val	11e ATC	ATT	Pro	8 6 F	Leu	GATG 1051	1171
Leu	สูย	Thr	Met	Ala GCC	Glu GAG	80 T CC	val GTT 70	Thr Leu Acc CTT 760	Ser Ser AGC AGC 850	GIU Ala GAA GCC 940	AAGG	GCTT
717 166	1rp V <sub>4</sub> 166 G	GIY Thr GGA ACA 220	Ser Met TCA ATG 310	Leu Ala CTG GCC 400	Ser Glu TCG GAG 490	Val Ser GTG TCC 580	ile val Att GfT 670		Ser AGC	GAA 9	GTCC 41	TACA 61
Het	Pro 7	Asn	Ser	<b>61.0</b>	Asp	GOA GOA	Ala	GLv GAA	Thr	Pro	ccre	AACT 11
ASP	Pro 1	Ile	Gly	Glu	Ser	Ale	Ala	gly gga	Pro	His	GTCT	CCAT
Asp	31n		11e ATA 300	G1y GGA 390	Ser TCT 480	Ser TCT 570	Val GTC 660	Met ATG	G1u GAG 840	Lys AAG 930	TAGT 1031	ACTC 1151
61y 668	Leu	150 166	Glu	Glu	Phe	Thr	Ala	GAA	Leu	Leu	GATC	AATG
Phe	Thr 1	Gln	Cys	Thr	GIn	Tyr	Ile ATT	Arg	G1y GGA	Leu	CCAG 21	TACT
Ser	6 U	. u.e		Leu CTC 80	Phe TTT	HIS Arg	Val Thr Gly Grc Acr GGG	E SAC	Ser	Ser TCA 920	10	11 11
Thr	val ile	Ser Thr TCC ACA	Glu Tyr Arg GAA TAC AGG	Val Leu GTC CTC 380	Ser Phe TCC TTT	HIS CAC	Thr	ASP GAT	Pro S CCA T	17r	CAGA	GTTG
Leu	Ala GCT	Ser	. ag	Arg	Lys	Arg Aga	Val	Pro	Pro	Thr	AAGC 1011	1131
Ile	8 % S	GAC	<u>&gt;</u> ;	Ser Arg	Arg Asn Gly Lys S	Thr Gly Arg	Ala Ala GCT GCT	53	Ser Pro Gly F AGC CCA GGG C	Ash The Ile The Tyr Ser AAC ACG ATC ACC TAC TCA 910	TGGCATTGGGAAAGCAAGCCAGAM 1001	tacgagattggttggttgactgtactaatgactccataacttacagcttcccaactcatctgctatcgatccactacactgccactaaattaatcaacttactgccgt 1121 1131 1131 1141 1151 1161 1171 1181 1191 1201 1211 1231
Met	Thr Lys Acc ARG	100 GIY ASP GGA GAC	Ser AGT	Ser	Ash	Thr	Ala	GLY	Pro	Thr	GGAA 01	TGG1
GAC	Ala 7	CCT	ASP	A P S	Arg	<b>в1</b> у 660	Val GTG	Pro	Ser	ABO	ATTG	iagat 11
Met Ile Leu Thr So TTT <u>STCGAG</u> ATG ATT CIT ACC AV	Asn A	te cro	Gla	Glu	TYE	Ser	Ile	Les CTC	val GTC	Glu	TGGC	TACG
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AGAGACTGCA

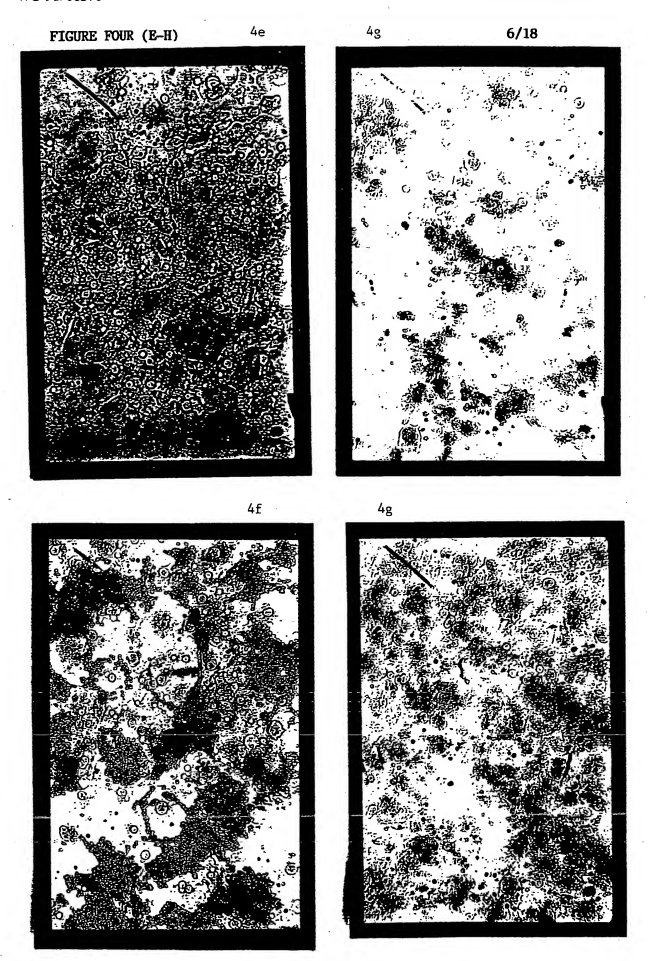
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FIGURE THREE B

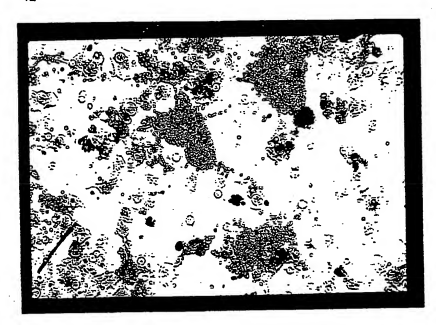
	- m -	0.71.00	L D m	m (h m	6 t) m	H (1 m			m11 m	11 (D. m.	~ m ~			
Met	val GTG 193	11e ATC 283	Ser AGC 373	Ar9 Agg 463	Phe TTC 553	Thr ACC 643	Cya TGT 733	Ser TCC 823	Arg CGC 913	Met ATG	Gln CAG 1093	GAA GAA 183	Thr ACT 273	
His CAT 100	Gln CAG	Ser	Leu CTC	Leu	Asn	val GTC	ABD	Thr	Lys	II. ATA	Glu GAG	Lys AAA 1	Gln CAA	
CYe	11e ATC	Arg Agg	Arg	Thr	Ser	Pro	Leu	Asn	Leu	G1y GGA	Ser	Pro	Ala	¥
Leu	a Ge	>0 -00	Thr ACC 163	IIIe ATC 53	e de la companya de l	Lye AAG 33	Thr Acc	Arg AGG	Val Grc 903	Val GTG		3 GA	G1y GGG 63	GAG
Thr L ACT T 90	Pro T CCG T	Asn G AAT GC 273	Gln 7 CAG A	Thr I	Tyr S TAC A	Ser 1 TCC A	Val T GTC A	Tyr A TAC A	or v 191 9	Ser V TCA G	Leu Val TrG GIT 1083	Thr T ACC A	Thr GACT G	Parice 373
	Pro Pro	His A CAC A	Glu G GRG C	Glu T GAA A	His T	Gin s CAG T	Leu V.	Glu T GAG T	Ser Ser AGC AG	Leu S CTG T	Pro L	Gln T	Ser Ti	1000
r Arg						9 D		១១ ១១	a U		a S S S			ATCC
e Ser	u Glu c GAG 173	p Phe G TTC 263	n Met A ATG 353	u Gly A GGG 443	Tyr Hie Tar Car 533	Leu His CIG CAC 623	y Ser G AGT 713	Ile Leu ATC CTG 803	Glu Asp GAG GAC 893	TYF T TAT 983	Glu Val GAA GTC 1073	S SOF G AGC 163	r Asp I GAC 253	3AGG 1363
Phe Trc 80	Leu CTC	Trp Trg	Glu I	5 8		35	61y 6 666 7	AT A	69	Phe TTT 98		Ala 6 666 116	AGT 125	CTA
Vel GTG	LYS	Gln	Cys	Leu	Arg	Thr	Glu	Lys	Thr	CTG	Leu	The	Pro	<b>GA</b> CT
His	val GTC	Thr	Arg CGG	Phe	Val	Arg Agg	Pro	Ser	Ala	116 ATC	A.n	Ala	Pro	36TC 135
Val GTC	val gtg 163	Ser TCT 253	Tyr Tat 343	Val GTG 433	Ser TCC 523	G1y GGA 613	Pho TTC 703	61y 66C 793	Val GTA 883	His CAC 973	TYE TAC 1063	Thr ACA 1153	Leu CTT	A SA
Thr ACT 70	Ala GCT	Ser	GAA	Leu	Ly. AAA	Leu	010	Val GTG	Glu GAG	Phe TTT	LYS AAA 1	Vel GTA	7. CCC 1	GGAJ 43
Trp	Lys AAG	ABIN	GLY	Gln	GLU	Ser	Ser	Tyr	Cys TGT	Trp	Ly.	GAA	GIn	GACT 13
Asn	Pro CCA 53	61y 666	Ser AGT 133	Pro CCT	Asn AAT 513	G1y GGA 603	Ser TCA 693	Phe TTC 183	Trp TGG	Val GTC	Glu GAG 53	Glu GAA	333	TACA
Ser AGC 60	CTT	Pro CCT	Asp GAC	Thr	HIS CAT	Lye	Val GTG	Ser	TAC	Pro CCT	AFG AGA 10	TYE TAT	Gln 1 CAG C	\ATA 1333
GAG 1	Asp ]	Asn I	Aen I	Gln 1 CAG 1	Phe I	Cys 1	Ser 1	Phe trrc	Phe 1	Ala E	Gln /	Val 1 Gre 1	Glu GAG	ctgtgggaccatggc <b>atctttaataaa</b> caaatatacagactggaaaaaggcgactctagaggatccccgg <u>dtggad</u> aaa 1303 1313 137 <u>3</u> 1373
Met G	AT	His P	TC P	Leu G CTC C	Phe F TTC 1	AC 1	Ala S GCA 1		GGA 1	Ser A	9 E	617 V 660 0	Pro G	TAA
	Thr H ACT C	Thr H ACC C 233	Thr Val ACA GTC 323	Leu L CTG C	Ser P TCG I	Tyr Tyr TAC TAC 593	AEG AAGA G	Leu His CTT CAC 773	Ala d GCT G 863	Ser S TCA T 953	Arg Leu AGG CTG 1043	Asp GAT GAT G	GLY P GGA C 1223	132
GGAATCCTGCCGTTCCTACTGATCCCC 30	>0	Gly T GGG A	Ala T GCC A	Leu L CTG C	Ile s ATC T	Asp T GAC T	Leu A CTG A	Gln L CAG C	Asp A GAT G	GIN S CAG T	His A	Ser A AGC G	CY. G TGT G	TCL
CTG	Ala GI GCT GG	Glu G GAA G	Lyb A.	Trp L TGG C	Arg I	61y A 666 6	Val Logical	TTA C	Glu A	Pro G	IJe H.	Arg Se Aga A	Asp C.	CAA 113
CCT														CATO
CGTT	u Ala T GCT 133	r Cys A 16C	r Phe G TTT 313	r Asp I GAC 403	B Asn G AAC 493	S Ser C AGT 583	r Pro	61y 666 763	A AGA B AGA B S 3	u Gly r GGT 943	Lys G AAA 1033	n Val A GTT 1123	1 Gly 3 GGT 1213	ATGG
CT GC	r CTT	Thr 3 ACA	Thr	S SOF	Leu A CTG	r His	Thr.	Pro	G GAA	Leu 3 CTT	r cre	CAA	r Val	3ACC.
30 30	AAT	Leu	TAC	110 ATT	Leu	Ser	Thr.	Arg AGA	Ala GCG	val GTG	TYE:	Glu	Ser	TGGC
	Leu CTA 123	Thr ACA 213	Ser AGC 303	Val GTG 393	Lys AAA 483	H15 CAC 573	Phe TTC 663	Gln CAG 753	Arg AGG 843	Gln CAA 933	Leu CTC 023	Phe TTT 113	Ser AGC 203	ACTG 3
FGAT	Val GTG	Val GTG	Ala GCC 3	G1y GGA	ABO	ABn AAC 5	រូ អូ សូ	CTG	A18 609	reu crc	Val 1 Grr (	Ser TCC 11	Arg CGA	IGAA 129
PAGC.	Ala	Thr	GP.	Leu	Arg	Ala	61y 666	CTC	ATA	G1u GAG	Thr	Aan	Pro	CCCTGAAA(
CCAC	rp Thr 3G ACA 113	Asp GAC	Val GTC 3	Asp GAT	Trp Tog	Lys AAA 3	GAA	Leu	H18 CAT	Leu TTG	Asn AAC 3	Ala GCA	GLY GGA 3	TGA
CGCI	Trp TGG	Glu Asp GAA GAC 203	Gln Val CAG GTC 293	Vel Asp GTA GAT 383	Ser Trp AGC TGG 473	Pro Lys CCC AAA 563	Val Gln GTC CAA 653	Asn Leu AAT TTG 743	Tyr His TAC CAT 833	Glu Leu GAG TTG 923	Leu Asn TTG AAC 1013	Lys Ala AAA GCA 1103	Asp Gly GAT GGA 1193	Ser *** AGT TGA 1283
GACT 1	CTG	Lys	Ser	Pro	HIS	ATC	Thr	Thr	GAG	Pro	Ser	Lys	OCC C	GAA
TGCAGACTCGCTCCAGAGCTGATG	CTA	Leu	Arg	Asp	Cys	Ser	ATC	GNG	Ser	NG C	Phe	GLY	A18 606	Ser

4c 43 5/18 FIGURE FOUR (A-D) 4d 4ь

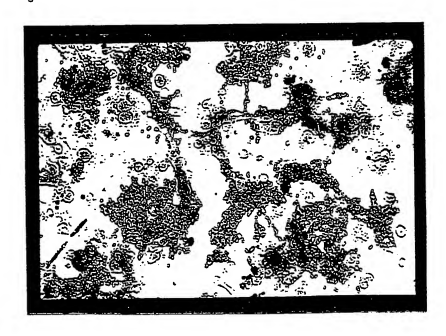
WO 91/06570 PCT/AU90/00513

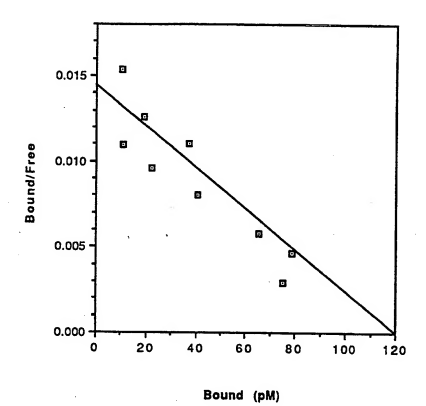


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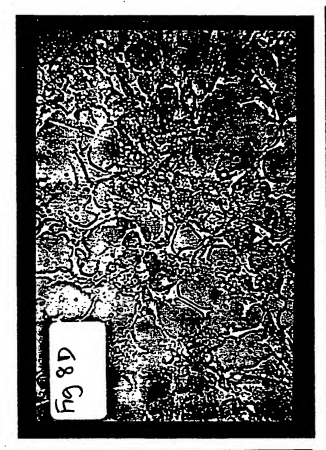
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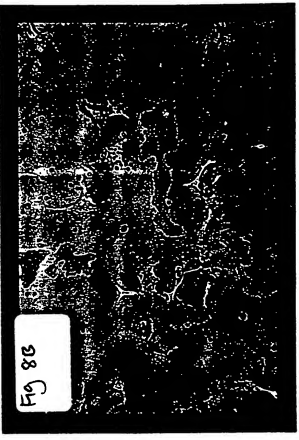




. 26	182	272	362	452	542	632	722	812	905	1002	1122 1242 1362 1370
Ala	Thr	Ser	Val GTG	Lys	His	Ser	Ile	118 ATC	Asp	FEC	CAN
Leu A	Val T GTG A	Pro S CCC A	Thr V ACT G		Asn H	2000	ودا	Ala j GCC J	Asp GAC	AGAGTAACGTTATGCCATGTGGTCATACTCTCAGCTACAGCTT	aatttaatggagactggaaaattcctgagcaaacaaaaccacctggcccttagaaataggacatgacctaacctaaacttaacttaacttaaactaaacca Faaccaaaaacttaactt
Leu L CTG C	Ser V TCT G	Gln P CAG C	Leu I	Lys A	Ala P GCA P	Val B	Ala 1 GCC 1	Ile /	The	AGCT	CCAA
Leu CTG CT	20 Asp S GAC T	50 Thr G ACG C	BO His L CAT C	110 Trp L TGG A	140 Gln A	GIn V		230 Met 1 ATG 7	260 Pro 1	icTC!	MATCA
Teau Tigo	Glu A	His T	Val H GTG C	Ser T AGC T	Pro G	Val G	Val V GTA G	Gin M	Ala P	TAC!	SACAT SACAT
Val Le GTT TO	Gln G	Thr H	Pro V	His S Cac a	Ile P ATC C	Thr V ACT G	Ala V GCT G	Arg G CGT C	Arg A	67.03	10001 11000 11000
Thr Ve ACA GI	Leu G	Pro II	Asp P	Cys H	Ser I	Ile T ATC A	Ala A GCT G	Gly A	Pro A	TOTO	CAAC
	Val Le GTG C!	Ile Pi ATT CO	Ser A	Arg C	Phe S TTC T	Thr I	Val A GTT G	Pro G CCT G	Asn P	TGCCA FG S	TAGA AAGA TATT
o Leu			Leu Se CTC AC	Leu An	Thr Pl	Val Ti GTG A	Ile V Att G	Pro P	Leu A	TTAT	CCCT
n Pro	Ile Asn ATC AAC		Ser Le	Met Le ATG CT	Pro 13	Pro CCT G	Ala I. GCC A	Glu P. GAG C	Thr L	AA CG	CTGG ATTG
u Gln T Caa		Gly Aen GGG AAT	Thr Se	Ile Me ATC A1	0.54	n ()	Ala A. GCA G	Phe G TTT G	Met I	GAGT.	CCAC
u Leu G CTT	o Trp		Gln Th	Thr II	TTG GA	3	Val A. GTA G	GIN P	Tyr M	*** TAA A	ANAA TGCT ACCA
20 rp Leu GG CTG	o Pro	s Asn		Glu Th Gaa ac	45	Ser Ser	Ala Ve GCT G1	A1A 6] GCC CJ	61y T	Asn *	AAAC GAAA
	o Pro	40 he His TC CAC	70 hr G1y CT GGC		o r Arg	1			250 G1y G1 GGC G(	280 Asn A	CTGG
r Leu C CTG	u Glu T GAG	₽ E	E 4	100 u Gly G GGA	130 e Ser	T 160 to Phe				Ser Ac	CCT VTTT VTGT
A AAC	e Leu	n Trp 3 TGG	e Gln c cAG	n Glu G GAG	s Phe	P L L	e Ala T GCG	1 Lys g Aag	A ASP		HATCH
Arg AGA	LY6	Gln r cAG	C TGV	G CAG	n Lys	r Thr	1 Ile C ATT	o Val T GTG	r Ala A GCT	1 Asn C AAC	TTGGA
Pro	Leu	ATT	Thr	Phe TTC	r Gln	1 24	1 Val	P Pro	u Thr A ACA	e Val	AGAC TAAC AAAA
Cys TGT	Val	Ser	TYE	Glu G GAG	Ser TCC	a G1y	a Val I GIG	r Asp r gar	r Glu T GAA	P His C CAT	ATGG AACT CCCC
Val	Ala	ASP GAC	Glu GAG	Leu	Lys	12 110 2 ATA	l Ala 3 GCT	r Thr	o Tyr	n Asp c gac	TTAA GCAA CTAC
Asn	Lys	Ser	61y 666	H18		EG 12	val r gre	n Ser r rcc	n Asp I GAC	o Asn C AAC	AAAT ATAA ATAC
g g	Pro	GAG	Ser	Pro	Gln Asn CAG AAT	Gly GGA	Ile ATT	Asn AAT	A ASD	P CCC	AGGA AAGC GAAG
Ser	<b>6</b> 50	Pro	Asp GAC	Thr Acc			7 Ile	A BCC	r Asn	u Pro r ccr	TTAN
-30 Met ATG	Ala GCT	Ser	Asn	Gln CAG	Phe	C T G T G T G T G T G T G T G T G T G T	61y 666	Ser	Thr A ACC	Leu CIT	OTAC
61n CAA	Ala GCA	30 Arg CGC	Asn AAC	Le 20	120 Phe TIC	150 His CAC	180 Met	210 I11e	240 Glu	270 Thr	SGGA
Thr	ALA GCT	Ala GCT	Asn	val GTG	val Thr	Tyr	Pro	Arg CGG	glu GAA	Leu	GAGG
Glu GAG	ola Ses	G13 GGG	Ala		Val GTC	asp gat	Ser	Lys	Leu	TYE	ACGO
-34 Met ATG	Ser	Gln	Lys	Trp 166	Lys	G1y GGT	Ser	Lys	Gla	Ile ATC	CTTC
ACT	A SA S	Cys	Phe	Glu GAA	Val	Ser AGT	Ser	Arg	Arg AGA	Asn	GATG
GAATTCCAACT	ALA GCA	Thr	Arg	Ser	Leu	H ts	Gly GGC	Cys TGC	Lys	Lys	GCTGAGTGGATGACAAAAAGGGGGAATTGTTAAAGGAAA ACACAAGCAAAACTTCACGGGGTCATACTACATACAAGCAT ACTAACATATAATTAGTGACTAGGGACTTTCTAAGAAGAT CTTGCCTG
GAAT	Ser	Leu	Tyr	Leu CTT	Pro	Ser	Met	Tyr	Arg Aga	Asp Gat	ACTG ACTA CTTA

FIGURE EIGHT (A-D)







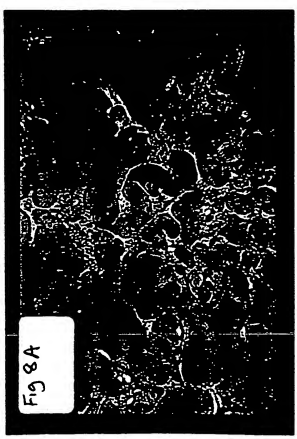


FIGURE EIGHT (E-H)

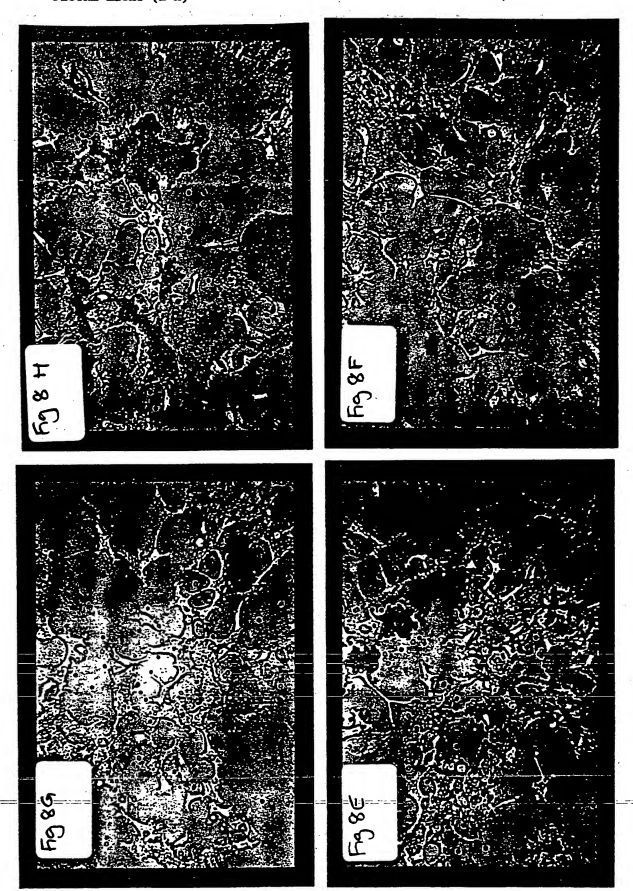


FIGURE EIGHT (I-J)



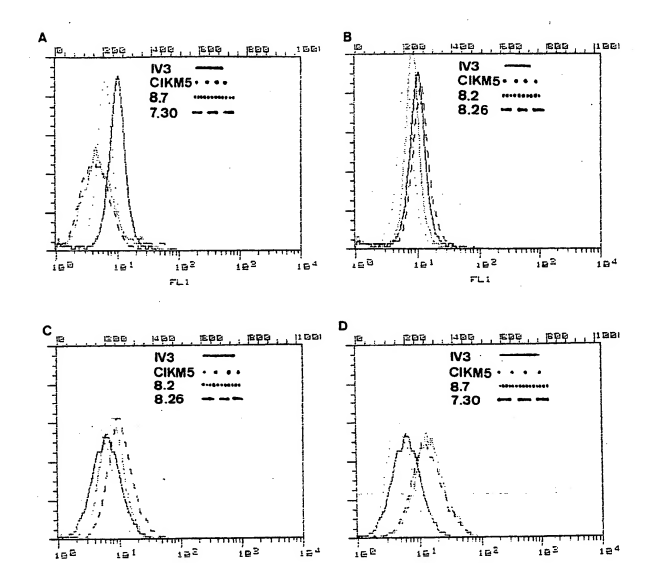


SUBSTITUTE SHEET

FIGURE NINE

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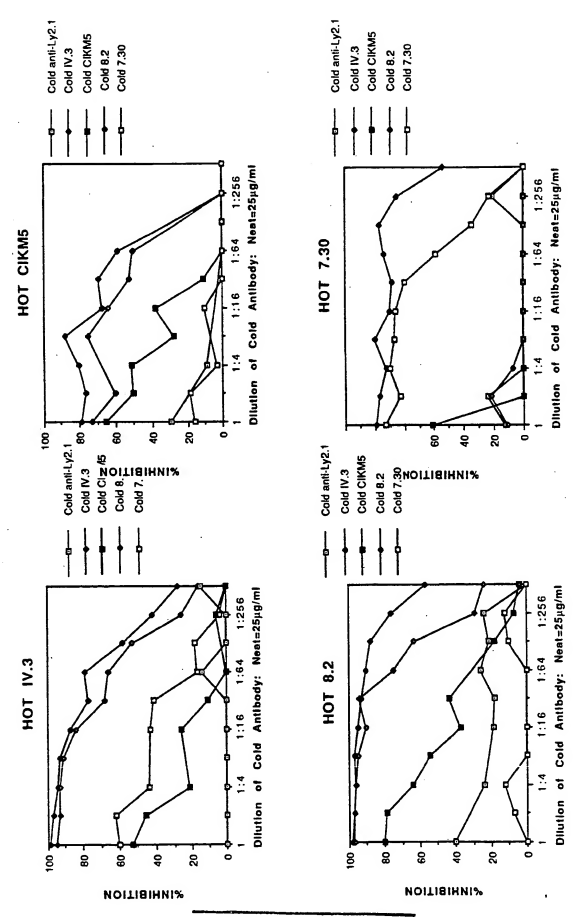
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FIGURE TEN



CHRCTITHTE CHEET

FIGURE ELEVEN A

17.10.90 Immunpptn NP-40 O/N Exposure 10% reducing

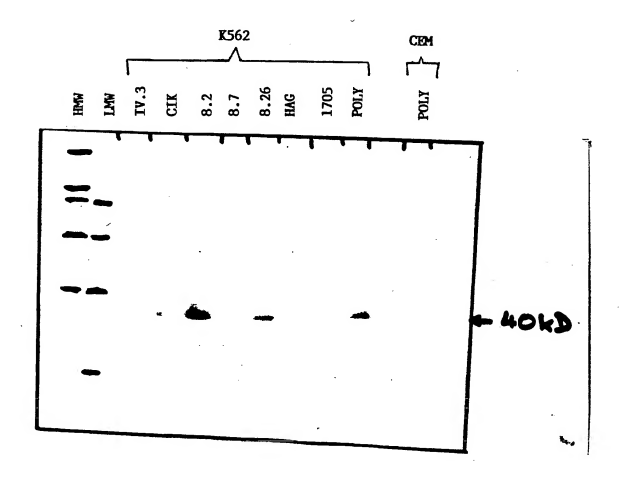
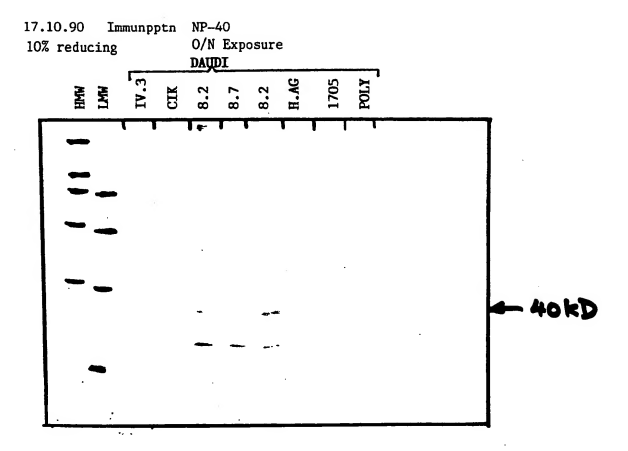
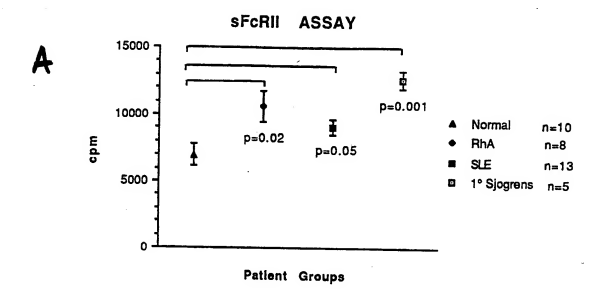


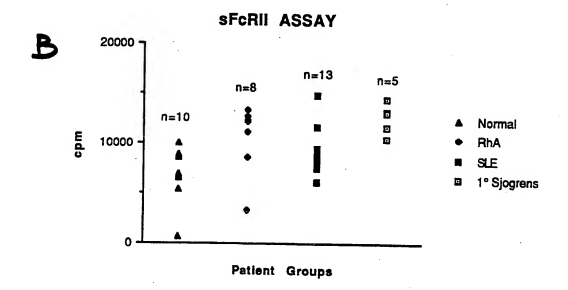
FIGURE ELEVEN B



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#### INTERNATIONAL SEARCH REPORT

International App (cation No. PCT/AU 90/00513

		The contact of the case of the				
I. CLA	SSIFICATION OF SURJECT MATTER (if several class	sification symbols apply, indicate all) 6				
According	g to International Patent Classification (IPC)	or to both National Classification and IPC				
Int. Cl.	5 C07K 15/06, 15/08, 15/12, C12N 15/13, C12P	21/08, GO1N 33/563				
II. FIE	LDS SEARCHED					
	Miniaum	Documentation Searched 7				
Classific	ation System   Classification	on Symbols				
IPC	WPAT : Keywords : "Immunoglo USPA : Keywords : As above	bulin", "Receptor", "Ig", "Fc", "FcR"				
	Documentation Searched other than Note to the Extent that such Documents are Include					
AU: CO7 Chemica	K 15/06, 15/08, 15/12, 13/00, CO7G 7/00, C12N 1 Abstracts : Keywords : "Immmoglobulin" or " "Fc Receptor" or "FcR	Ig" and "Binding Molecule";				
III. DOC	LIMENTS CONSIDERED TO BE RELEVANT 9					
Category*	Citation of Document, with indication, of the relevant passages					
A	AU,A, 32842/89 (SCHERING BIOTECH CORPORATION	) 5 July 1989 (05.07.89)				
<b>A</b> .	AU,A, 27830/88 (KISHIMOTO, T.) 13 July 1989	(13.07.89)				
<b>A</b>	AU,A, 82701/87 (MEMORIAL SLOAN-KETTERING CANCER INSTITUTE) 23 June 1988 (23.06.88)					
A	AU,A, 74881/87 (THE UNIVERSITY OF MELBOURNE) (04.02.88)	4 February 1988				
A	AU,A, 63858/86 (MEDICAL BIOLOGY INSTITUTE) 1	4 April 1988 (14.04.88)				
<b>A</b>	EP,A, 269455 (TAKEDA CHEMICAL INDUSTRIES, LTD) 1 June 1988 (01.06.88) (continued)					
* Spe	ecial categories of cited documents: 10 *T*	later document published after the				
art	cument defining the general state of the twhich is not considered to be of rticular relevance	international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention				
	rlier document but published on or "X"	document of particular relevance; the claimed invention cannot be considered novel				
"L" doc cla pui						
*0* do	cument referring to an oral disclosure,	involve an inventive step when the document				
"P" do	use, exhibition or other means is combined with one or more other such document published prior to the documents, such combination being obvious international filing date but later than a person skilled in the art.					
the	e priority date claimed "&"	document member of the same patent family				
IV. CE	RTIFICATION					
•	the Actual Completion of the	Date of Mailing of this International				
	ional Search	Search Report				
i	ional Searching Authority					
Australi	ian Patent Office	Lea Lia K. AYERS				

FURTHER I	NFORMATION CONTINUED FROM THE SECOND SHEET	
A	Hibbs, M.L. et al. Proc. Natl. Acad. Sci. USA, Volume 85, issued   April 1988, "Molecular cloning of a human immunoglobulin G Fc   receptor" see pages 2240-2244	
A	Shimizu, A. et al. Proc. Natl. Adad. Sci. USA, Volume 85, issued   March 1988, "Human and rat mast cell high-affinity immunoglobulin   E receptors: Characterization of putative ~ -chain gene products"   see pages 1907-1911	
P,A	Allen, J.M. and B. Seed. Science, Volume 243, issued 20 January 1989, "Isolation and expression of functional high-affinity Fc receptor complementary DNAs" see pages 378-381	·
A	Hibbs, M.L. et al. Proc. Natl. Acad. Sci. USA, Volume 83, issued   September 1986, "The murine Fc receptor for immmoglobulin:   Purification, partial amino acid sequence, and isolation of cDNA   clones" see pages 6980-6984	
V. []	OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1	

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

- 1.[] Claim numbers ..., because they relate to subject matter not required to be searched by this Authority, namely:
- 2.[] Claim numbers , because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
- 3.[] Claim numbers ..., because they are dependent claims and are not drafted in accordance with the second and third sentences of PCT Rule 6.4 (a):

#### VI. [ ] OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 2

This International Searching Authority found multiple inventions in this international application as follows:

- 1.[] As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims of the international application.
- 2.[] As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims:
- 3.[] No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:
- 4. [] As all searchable claims could be searched without effort justifying an additional fee, the International Searching Authority did not invite payment of any additional fee.

Remark on Protest

- [ ] The additional search fees were accompanied by applicant's protest.
- [ ] No protest accompanied the payment of additional search fees.

# ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL APPLICATION NO. PCT/AU 90/00513

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Document Cited in Search Report			Patent Family Members					
AU	32842/89	EP	319307	WO	8905351			
AU	27380/88	DK HU NO ZA	7124/88 50498 885687 8809506	EP IL PT	321842 88743 89299	FI JP EP	885873 2000443 321601	
AU	82701/87	WO	8803172				·	
AU	74881/87	EP	268636	JP	63503386	WO	8707277	
EP	269455	JP	63246398					

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